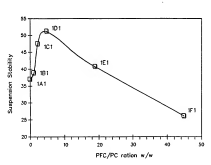


# WORLD INTELLECTUAL PROPERTY ORGANIZATION



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#### (57) Abstract

Engineered particles are provided for the delivery of a blosetive agent to the respiratory tract of a patient. The particles may be used in the form of any powders or in the form of stabilized dispensions comprising a nonequeous continuous phase. In particularly preferred mobidiments the particles may be used in conjunction with an inhaltant over excell as a day powder inhalter, metered dose inhalter or a nebulizer.

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#### PERFORATED MICROPARTICLES AND METHODS OF USE

#### Field of the Invention

The present invention relates to formulations and mathods for the production of perforated microstructures which comprise an active agent. In particularly preferred embodiments, the active agent will comprise a biosocive agent. The perforated microstructures will preferably be used in conjunction with inhalation devices such as a meterad does inhaler, dry powder inhaler or nebutizer for both topical and systemic delivery via pulmonary or nest routes.

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### Background of the Invention

Targeted drug delivery means are particularly desirable where toxicity or biosvaliability of the pharmaceutrical compound is an issue. Specific drug delivery methods and compositions that effectively deposit the compound at the site of action potentially serve to minimize toxic side effects, lower dosing requirements and decrease therapeutic costs. In this regard, the development of such systems for pulmonary drug delivery has long been a good of the pharmaceutical industry.

The three most common systems presently used to deliver drugs locally to the pulmonary eir passages are dry powder inhalters (IPFlat, metered dose inhalters (MDIs) and nebulizers. MDIs, the most popular method of inhaltetion administration, may be used to deliver medicaments in a solubilized form or as dispersion. Typic-lally MDIs comprise a Freon or other relatively high vepor pressure propelant that forces sensodized medication into the respiratory tract upon activation of the device. Unlike MDIs, DPIs generally rely entirely on the poeint's inapitatory efforts to introduce a medicament in a dry powder form to the lungs. Finally, nebulizers form a medicament sensul to be inhaled by imperting energy to a liquid solution. More recently, direct pulmonary delivery of drugs during liquid ventilation or pulmonary leavegu using a fluorochemical medium has also been explored. While each of these methods and associated systems may prove effective in selected situations, inherent drawbacks, including formulation limitations.com limit their ince.

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The MOI is dependent on the propulsive force of the propellant system used in its manufacture. Traditionally, the propellant system has consisted of a mixture of chlorofluorocarbons (CFCs) which are selected to provide the desired vapor pressure and suspension stability. Currently, CFCs such as Freen 11, Freen 12, and Freen 114 ere the most widely used propellants in aerosol formulations for inhalation administration. While such systems may be used to deliver solubilized drup, the selected bioscrive agent is typically incorporated in the form of a fine periculate to provide a dispersion. To minimize or prevent the problem of aggregation in such systems, surfactants are often used to coat the surfaces of the bioscrive agent and assist in wetting the particles with the senses propellant. That use of surfactants in this way to maintain substantially uniform dispersions is said to "Stabilize" the suspensions.

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Unfortunately, usditional chlorofluorocerbon propellents are now believed to deplete stratespharic ozons and, as a consequence, are being phased out. This, in turn, has led to the development of seroed formutations for pulmonary drug delivery employing so-called envirormentally friendly propellents. Classes of propellents which are believed to have minimal atons-depletion potential in companion with CPCs are perfluorinated compounds (PPCs) and hydrofluorosekmens (IPFAs). While selected compounds in these classes may function effectively as biocompatible propellents, many of the surfactunts that were effective in stabilizing drug suspensions in CPCs are no longer effective in these new propellent systems. As the solubility of the surfactant in the IFA decreases, diffusion of the surfactant to the interface between the drug particle and IFFA becomes exceedingly slow, leading to poor wetting of the medicament particles and a loss of suspension stability. This decreased solubility for surfactants in IFFA propellents is likely to result in decreased efficiency with regard to any incorporated bioactive agent.

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More generally, drug suspensions in liquid fluorochemicals, including HFAs, comprise heterogeneous systems which usually require medispersion prior to use. Yet, because of factors such as patient compliance obtaining a relatively homogeneous distribution of the phermaceutical compound is not always easy successful. In addition, prior art formulations comprising micronized perticulates may be prene to aggregation of the particles which can result in inadequate delivery of the drug. Crystal growth of the suspensions via Ostwald ripensing may also lead to particle size haterogeneity and can significantly reduce the shelf-life of the formulation. Another problem with conventional dispersions comprising micronized dispersants is particle coatsening. Coarsening may occur via several mechanisms such as flocculation, fution, molecular diffusion, and coalescence. Over a relatively short period of time these processes can cause the formulation to the point, where it is no loager usable. As such, while conventional systems comprising fluorochamical suspensions for MDIs or liquid ventilation are certainly a substantial improvement over prior art non-fluorochamical delivery vahicles, the drug suspensions may be improved upon to anable formulations with improved stability that also offer more efficient and accurate dosing at the desired size.

Similarly, conventional powdered preparations for use in DPIs often fail to provide accurate, reproducible dosing over extended periods. In this respect, those skilled in the art will appreciate that conventional powders (i.e. micronized) tend to aggregate due to hydrophobic or electrostatic interactions between the fine particles. These changes in particle size and increases in cohesive forces over time tend to provide powders that give undesirable pulmonary distribution profiles upon activation of the device. More particularly, fine particle aggregation disrupts the serodynamic properties of the powder, thereby preventing large amounts of the senselized medicament from reaching the deeper airways of the lung where it is most effective.

In order to overcome the unwanted increases in cohesive forces, prior art formulations have typically used large cernier particles comprising lectose to prevent the fine drug particles from aggregating. Such carrier systems allow for at least some of the drug particles to lossely bind to the lactose surface and

disengage upon inhibition. However, substantial amounts of the drug fail to disengage from the large factoss particles and see deposited in the throat. As such, these carrier systems are relatively inefficient with respect to the fine particle fraction provided per actuation of the DPI. Another solution to particle aggregation is proposed in WD 98/31348 wherein particles having relatively large geometric diameters (i.e. prefarably greater than 10 Jml) are used to reduce the amount of particle interactions thereby preserving the flowability of the powder. As with the prior art carrier systems, the use of large particles apparently reduces the overall surface area of the powder preparation reportedly resulting in improvements in flowability and fine particle fraction. Unfortunately, the use of relatively large particles may result in dozing limitations when used in stendard DPIs and provide for less than optimal dozing due to the potentially prelonged dissolution times. As such, there still remains a need for standard sized particles that resist aggregation and preserve the flowability and dispersibility of the resultine provider.

 Accordingly, it is an object of the present invention to provide mathods and preparations that advantageously allow for the nesal or pulmonary administration of providers having relatively high fine particle fractions.

It is a further object of the present invention to provide stabilized preparations suitable for serosolization and subsequent administration to the pulmonary air passages of a patient in nead thereof.

It is yet another object of the present invention to provide powders that may be used to provide stabilized dispersions.

It is still a further object of the present invention to provide powders exhibiting relatively low cohesive forces that are compatible for use in dry powder inhalers.

#### Summary of the Invention

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These and other objects are provided for by the invention disclosed and claimed herriin. To that and, the methods and associated compositions of the present invention previde, in a broad aspect, for the improved delivery of agents to a desired site. More particularly, the present invention may provide for the delivery of bioactive agents are sincered physiological target sites using perforated microstructure providers. In preferred embodiments, the bioactive agents are in a form for administration to at least a portion of the putmonary air passages of a patient in need thereof. To that end, the present invention provides for the formation and use of perforated microstructures and delivery systems comprising such providers, as well as individual components thereof. The disclosed providers may further be dispersed in selected suspension media to provide stabilized dispersions. Unlike prior and poveders or dispersions for drug delivery, the present invention preferrably employs novel techniques to reduce attractive forces between the particles. As such, the disclosed providers exhibit improved flovability and dispersibility while the disclosed dispersions exhibit reduced degradation perferably comprise a suspension medium e.g., a fluorechemical that further serves to reduce the rate of thermidation with mesure to the incomponents between the accordinate, the

dispersions or powders of the present invention may be used in conjunction with metered dose inhalers, dry powder inhalers atomizers, nebulizers or liquid dose instillation (LDI) techniques to provide for affective drug delivery.

With regard to particularly preferred embodiments, the hallow endlor porous perforated microstructures substantially reduce attractive molecular forces, such as van der Weals forces, which dominate prior art powdered preparations and dispersions. In this respect, the powdered compositions typically have relatively low bulk dominate which contribute to the flowability of the preparations while providing the desired cheracteristics for inhibitation therepies. More particularly, the use of multively low density perforated for porousl microstructures or microparticulates significantly reduces attractive forces between the particles thereby lowering the shear forces and increasing the flowability of the resulting powders. The relatively low density of the perforated microstructures also provides for superior serodynamic perforance when used in inhabitation therapy. When used in dispersions, the physical characteristics of the powders provide for the formation of stable preparations. Moreover, by selecting dispersion in accordance with the teachings benefit, interparticle attractive forces may further be reduced to provide formide formulations having enhanced stability.

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Accordingly, select embodiments of the invention provide for powders having increased dispersibility comprising a plurality of perforated microstructures having a bulk density of less than about 0.5 g/cm<sup>2</sup> wherein said perforated microstructure powder comprises an active agent.

With regard to the perforated microstructures, those skilled in the art will appreciate that they may be formed of any biocompatible material providing the desired physical characteristics or morphology. In this respect, the perforated microstructures will preferably comprise pores, voids, defects or other interestial spaces that can reduce attractive forces by minimizing surface interactions and decreasing sheer forces. Yet, given these constraints, it will be appreciated that any material or configuration may be used to form the microstructure matrix. As to the selected materials, it is desirable that the microstructure incorporates at least one surfactant. Preferably, this surfactant will comprise a phospholipid or other surfactant approved for pulmonary use. Similarly, it is praferred that the microstructures incorporate at least one active agent which may be a bioactive agent. As to the configuration, particularly preferred embodiments of the invention incorporate sprey dried, believe microsphares having a relatively thin prous wall defining a large internal void, although, other void containing or perforated structures are contemplated as well. In preferred embodiments the perforated microstructures will further comprise abiascrive expent.

Accordingly, the present invention provides for the use of a bioactive agent in the manufacture of a medicament for pulmonary delivery whereby the medicament comprises a plurality of perforated microstructures which are sensotized using an inhalation device to provide sensotized medicament comprising said bioactive agent wherein said accordized medicament is administered to at least a portion of the resid or pulmonary air passages of a patient in need thereof.

It will further be appreciated that, in selected embodiments, the present invention comprises methods for forming perforated microstructures that exhibit improved dispersibility. In this repard, it will be appreciated that the

disclosed perforated microstructures reduce attractive indecidar forces, such as van der Waals forces, which dominate prior art providend preparations. That is, unalke prior art preparations comprising relatively dense, solid particles or nonporous particles larg, microsized, the powdered compositions of the present invention exhibit increased flowobility and dispensibility due to the lower shear forces. In part, this reduction in cohesiva forces is a result of the novel production methods used to provide the desired providers.

As such, preferred embodiments of the invention provide methods for forming a perforated

providing a liquid feed stock comprising an active agent;

atomizing said liquid feed stock to produce dispersed liquid droplets;

drying said liquid droplets undar predetermined conditions to form perforated microstructures comprising said active agent: and

collecting said perforated microstructures.

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With regard to the formation of the perforated microstructures it will be appreciated that, in preferred embodiments, the particles will be spray dried using commencially available equipment. In this regard the feed stock will perfeasibly comprise a blowing agent that may be adected from fluorinated compounds and nonfluorinated oils. Preferably, the fluorinated compounds will have a boiling point of greater than about 80°C. Within the context of the instant invention the fluorinated blowing agent may be retained in the perforated microstructures to further increase the dispersibility of the resulting powder or improve the stability of dispersions incorporating the same. Further, nonfluorinated oils may be used to increase athe solubility of seventee bioscrive agents is up. steroids in the feed stock, resultion in increased concentrations of bioscrive expents in the perforated microstructures.

As discussed above, the dispersibility of the perforated minostructure powders may be increased by reducing, or minimizing, the van der Waals attractive forces between the constituent perforated microstructures. In this regard, the present invention further provides methods for increasing the dispersibility of a powder comprising the steps of:

providing a liquid feed stock comprising an active agent; and

spray dying said Kajud feed stock to produce a performed minostructure powder having a built density of less then about 0.5 plcm<sup>2</sup> wherein said powder exhibits reduced van der Waals attractive forces when compared to a relatively non-porous powder of the same comparision. In particularly preferred embodiments the perforated microstructures will comprise hollow, prous microspheres.

The blowing agent may be dispersed in the carrier using techniques known in the art for the production of homogenous dispersions such a sonication, mechanical mixing or high pressure homogenization. Other methods contemplated for the dispersion of blowing agents in the feed solution include co-mixing of two fluids prior to atomization as described for double nebulization techniques. Of course, it will be appraiciated that the atomizer can be customized to optimize the desired particle characteristics such as particle size. In special cases a double liquid nazzle may be employed. In another embodiment, the blowing

agant may be disparsed by introducing the agent into the solution under elevated pressures such as in the case of nitrogen or cerbon dioxide gas.

As to the delivery of perforated microstructure powders or stabilized dispersions, another aspact of the prasant invention is directed to inhelation systems for the administration of one or more bioactive agents to a patient. As such, the present invention provides systems for the pulmonary administration of a bioactive agent to a patient comprising:

an inhalation device comprising a reservoir: and

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a powder in said reservoir wherein said powder comprises a plurality of perforated microstructures having a bulk density of less than about 0.5 glcm² wherein said perforated microstructure powder comprises a bioactive agent whereby said inhalation device provides for the serosatized administration of said powder to at least a portion of the pulmonary air passes of a patient in need thereof. As alluded to above, it will be appreciated that an inhalation device may comprise an atomizer, a sprayer, a dry powder inhaler, a matered doss inhaler or a nebulizer. Moreover, the reservior may be a unit dose container or bulk reservior.

In other emodiments, the perforated microstructure powders may be dispersed in an appropriate suspension medium to provide stabilized dispersions for delivery of a selected agent. Such dispersions are particularly useful in metered dose inhalers and nebulizers. In this regard, particularly preferred suspensions mediums comprise fluorochemicals (e.g., particurocarbons or fluorocebroids) that are liquid at more temperature. As discussed above, it is well established that many fluorochemicals have a proven history of safety and biocompatibility in the lurg-Further, in contrast to aqueous solutions, fluorochemicals do not regatively impact gas acchange. Moraover, because of their unique wettability characteristics, fluorochemicals may be able to provide for the dispersion of particles desper into the lung, thereby improving systemic delivery. Finally, many fluorochemicals are also bacteriotateic thereby decreasing the potential for microbal growth in composible preparations.

Whether administrated in the form of a dry powder or stabilized dispersion, the present invention provides for the effective delivery of bioactive agents. As used herain, the terms "bioactive agent" refers to a substance which is used in connection with an application that is therapeutic or diagnostic in thems. such as methods for diagnostic plane presents or absence of a disease in a patient and/or methods for treating disease in a patient. As to correstible bioactive agents, those skilled in the art will appreciate that any therapeutic or diagnostic agent may be incorporated in the stabilized dispursions of the present invention. For exemple, the bioactive agent may be selected from the group consisting of antiallergics, bronchodistors, bronchodistors, pulmonary lung surfactants, analgeaics, antibiotics, leukotrione inhibitors or antagonists, anticholinergics, mast cell inhibitors, antibiotars antifical manatories, antihopicalistics, ansi-tuberculars, imaging agents, or disconstitute agents, anomes, steroids, genetic meterial, wire vectors, emissengents, proteins, peptides and combinistors thereof. In preference anobdiments the bioactive agents comprises composeds which are to be administrated systemically fize, to the systemic circulation of a patient guch as peptides, pretrains or polynucleoticies. As will be discossed in more detail

below, the bioactive agent may be incorporated, blended in, coated on or otherwise associated with the perforated microstructure.

Accordingly, the present invention provides methods for the pulmonary delivery of one or more bioactive epents comprising the steps of:

providing a powder comprising a plurality of perforated microstructures having a bulk density of less
than about 0.5 g/cm² wherein said perforated microstructure powder comprises a bioactive agent;

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aerosolizing said perforated microstructure provider to provide an aerosolized medicament; and administaning a therapeutically effective amount of said aerosolized medicament to at least a portion of the nasei or pulmonery passages of a patient in need thereof,

As used herein tha term "enrosolized" shell be held to mean a gessous suspension of fine solid or liquid particles unless otherwise dictated by contextual restraints. That is, an aerosol or aerosolized madicament may be generated, for example, by a dry powder inhalar, a matered dose inhalar, an atomizer or a nebulizar.

With respect to the disclosed powders, the selected agent or bioactive agent, or agents, may be used as the sole structural component of the perforated microstructures. Conversely, the perforated microstructures may comprise one or more components (i.e. structural materials, surfactants, excipients, etc.) in addition to the incorporated agent. In particularly prefarred embodements, the suspended perforated microstructures will comprise relatively high concentrations of surfactant (greater than about 10% which along with an incorporated bioactive agent(s). Finally, it should be appreciated that the particulate or perforated microstructure may be costad, linked or between associated with an agent or bioactive agent in a non-integral manner. Whatever configuration is selected, it will be appreciated that any associated bioactive agent may be used in its natural form, or as one or more salts incovers in the act.

While the providers or stabilized dispersions of the present invention are particularly suitable for the pulmonary signification of bioactive agents, they may also be used for the localized or systemic administration of compounds to any location of the body. Accordingly, it should be emphasized that, in preferred embodiments, the formulations may be administrated using a number of different routes including, but not limited to, the gastrointestinel tract, the respiratory tract, topically, intramuscularly, intrapentoneally, neaally, vaginally, acctally, surally or coularly.

Other objects, features and advantages of the present invention will be apparent to those skilled in the art from a consideration of the following detailed description of preferred exemplary embodiments thereof.

## Brief Description of the Drawings

Figs. 1A1 to 1F2 illustrate changes in particle morphology as a function of varietion in the ratio of fluorocarbon blowing agent to phospholipid (PFCIPC) present in the sprey dry feed. The micrographs, produced using scanning electron microscopy and transmission electron microscopy techniques, show that in

the absence of FCs, or at low PFC/PC ratios, the resulting spray dried microstructures comprising gentamicin sulfate are neither particularly hollow nor porous. Conversely, at high PFC/PC ratios, the particles contain numerous pores and are substantially hollow with thin wells.

Fig. 2 depicts the suspension stability of gentanicin particles in Perflubron as a function of formulation PFC/PC ratio or particle porosity. The particle porosity increased with increasing PFC/PC ratio. Maximum Stability was observed with PFC/PC ratios between 3 to 15, illustrating a preferred morphology for the northborn suspension media.

Fig. 3 is a scanning electron microscopy image of perforated microstructures comprising cromolyn sodium illustrating a preferred hollow/porous morphology.

Figs. 4A to 4D are photographs illustrating the enhanced stability provided by the dispersions of the present invention over time as compared to a commercial cromolyn sodium formulation lintal", Rhone-Powent-Roserl. In the photographs, the commercial formulation on the left rapidly separates while the dispersion on the right, formed in accordance with the teachings herein, remains stable over an extended period.

Fig. 5 presents results of in-vitro Andersen cascade impactor studies comparing the same hollow prous albutard sulfate formulation delivered via a MOI in HFA-134a, or from an exemplary DPI. Efficient delivery of particles was observed from both devices. MOI delivery of the particles was maximized on plate 4 corresponding to upper airway delivery. DPI delivery of the particles results in substantial deposition on the later states in the impactor corresponding to improved systemic delivery in-vivo.

#### Detailed Description Preferred Embodiments

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While the present invention may be embodied in many different forms, disclosed herein are specific illustrative embodiments thereof that exemplify the principles of the invention. It should be emphasized that the present invention is not limited to the specific embodiments illustrated.

As discussed above, the present invention provides methods, systems and compositions that comprise perforated microstructures which, in praferred embodiments, may advantageously be used for the delivery of bioactivo agents. In particularly preferred embodiments, the disclosed perforated microstructure providers may be used in a dry state (e.g., as in a DFI) or in the form of a stabilized dispersion (e.g., as in a MDI, LIO or nebulizer formulation) to deliver bioactive agents to the nasal or pulmonary air passages of a patient. It will be appreciated that the perforated microstructures disclosed herein comprise a structural metrix that exhibits, defines or comprises voids, pores, defects, hellows, spaces, interstitial spaces, apertures, perforations or holes. The elabolute shape (as opposed to the morphology) of the perforated microstructure is generally not critical and any overall configuration that provides the desired characteristics is contamplated as being within the scope of the invention. Accordingly, preferred embodiments can comprise approximately microspherical shapes. However, collapsed, deformed or fractured periodiates are also compatible. With this caveat, it will further is approximate that, periodically preferred embodiments of the invention comprise gray died hollow, prorus microspheres. In any

case the disclosed powders of perforeted microstructures provide several adventages including, but not limited to, increases in suspension stability, improved dispersibility, superior sampling characteristics, elimination of carrier perticles and enhanced serodynamics.

These skilled in the art will apprecise that many of these aspects are of perticular use for dry powder inhaler applications. Unlike prior art formulations, the present invention provides unique methods and compositions to reduce cohesive forces between dry particles, thereby minimizing particulate eggregation which can result in an improved delivery efficiency. As such, the disclosed preparations provide a highly flowshile, dry powders that can be efficiently excessioned, uniformly delivered and penetrate deeply in the lung or nesal pesseges. Furthermore, the perforced microstructures of the present invention result in surprisingly low throat decostion uson administration.

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In preferred embodiments, the perforated microstructure powders have relatively low bulk density, ellowing the powders to provide superior sampling properties over compositions known in the ert. Currently, as explained ebove, many commercial dry powder formulations comprise large lactose perticles which have micronized drug aggregated on their surface. For these prior art formulations, the lactose particles serve as a carrier for the active agents end as a bulking agent, thereby providing means to partiely control the fine particle doss delivered from the device. In addition, the lactose particles provide the means for the commercial filling capability of dry particles into unit dose containers by adding mess and volume to the desage form.

By way of contrast, the present invention uses methods and compositions that yield powder formulations having extraordinanily law bulk density, thereby reducing the minimal filling weight that is commercially feasible for use in dry powder inhaletion divices. That is, most unit does containers designed for DPIs are filled using fixed volume or grevimetric techniques. Contrary to prior art formulations, the repeated invention provides powders wherein the active or bioactive agent and the incipients or bulking agents make up the entire inhaled perticle. Compositions according to the present invention typically yield powders with bulk densities less than 0.5 glam<sup>2</sup> or 0.3 glam<sup>2</sup>, preferebly less 0.1 glam<sup>2</sup> and most preferebly less than 0.05 glam<sup>3</sup>. By providing particles with very low bulk density, the minimum powder mess that can be filled into a unit does container is reduced, which eliminates the need for carrier particles. That is, the relatively low does phemaceutical compounds. Moreover, the elimination of carrier particles will potentially minimize throat deposition and eny "gag" effect, since the large lactase particles will impact the threat and upper sirvexes due to their size.

In accordance with the teachings herein the perforeted microstructures will preferably be provided in e "dry" state. That is the microparticles will possess a moisture content that allows the powder to remain chemically and physically stable during storage at ambient temperature and easily dispersible. As such, the moisture content of the microparticles is typically less than 6% by weight, and perferably less 3% by weight.

In some instances the moisture content will be as low as 1% by weight. Of course it will be appreciated that the moisture content is, at least in part, dicteted by the formulation and is controlled by the process conditions employed, e.g., inlet temperature, feed concentration, pump rate, and blowing egent type, concentration and test divino.

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With respect to the composition of the structural matrix defining the perforated microstructures, they may be formed of any material which possesses physical and chemical characteristics that are compatible with any incorporated active agents. While a wide variety of materials may be used to form the particles, in particles, preferred phermaceutical embodiments the structural matrix is associated winh, or comprises, e surfectant such as phaspholigio of fluorinated surfactent. Although not required, the incorporation of a compatible surfactant cen improve powder fluoveity, increase aerocal efficiency, improve dispersion stability, and facilitate preparation of a suspension. It will be appreciated that, as used herein, the terms "structural matrix" or "nicrostructure matrix" are equivalent and shall be held to mean any solid material forming the perforated microstructures which define a plurality of voids, apertures, hollows, defects, ponss, holes, fissures, etc. that provide the desired characteristics. In preferred embodiments, the perforated microstructure defined by the structural matrix components are provided in the proper companies a spray dised hollow porous microsphere incorporating at least one surfactant. It will further be operaccampated that, by altering the matrix components, the density of the structural matrix may be edjusted. Finally, as will be discussed in turther detail below, the perforated microstructures perforably compass at least one solve or bisactive agent.

As indicated, the perforeted microstructures of the present invention may optionally be associated with, or comprise, one or more surfactants. Moreover, miscable surfactants may optionally be combined in the case where the microparticles are formulated in a suspension medium liquid phase. It will be appreciated by those skilled in the art that the use of surfactants, while not necessary to practice the instant invention, may further increase dispersion stability, provider flowability, simplify formulation procedures or increase efficiency of delivery. Of course combinations of surfactants, including the use of one or more in the liquid phase and one or more essociated with the perforated microstructures are contemplated as being within the scope of the invention. By "associated with or comprise" it is meent that the structural matrix or perforated microstructure may incorporate, adsorb, absorb, be comed with or a formed by the sufficiency.

In a broad sense, surfactants suitable for use in the present invention include any compound or composition that eids in the formetion of perforated microparticles or provides enhanced suspension stability, improved powder dispersibility or decreased particle aggregation. The surfactant may compisse a single compound or any combination of compounds, such as in the case of co-surfactants. Particularly preferred surfactants are nonfluorinated and selected from the group consisting of saturated and unsaturated lipids, nonionic detargents, nonionic block copolymers, ionic surfactants and combinations thereof. In those embodiments comprising stabilized dispersions, such nonfluorinated surfactants will preferrably be relatively insoluble in the suspension madium. It should be emphasized that, in addition to the efferementioned surfactants, suitable fluorinated surfactants are compatible with the teachings herein and may be used to provide the desired preparations.

Ligids, including phospholipids, from both natural and synthetic sources are particularly compatible with the present invention and may be used in varying concentrations to form the structural matrix. Generally compatible lipids comprise those that have a gel to liquid crystal phase transition greater than about 40°C. Preferably the incorporated lipids are relatively long chain (i.e. C1x C2) seturated lipids and more preferably comprise phospholipids. Exemplary phospholipids useful in the disclosed stabilized preparations comprise, dipalmitovlphosphatidylcholine, disterpylphosphatidylcholine, diarachidoylphosphatidylcholine dibehennylnhosnhatidylcholine. short-chain nhosphatidylcholines. long-chain saturated phosphatidylethanolamines, long-chain saturated phosphatidylserines. Iong-chain saturated phosphatidylalycerols, long-chain saturated phosphatidylinositols, glycolipids, ganglioside GM1, sphingomyelin, phosphatidic acid, cardiolipin; lipids bearing polymer chains such as polyethylene glycol, chitin, hyaluronic acid, or polyvinylpyrrolidona; lipids bearing sulfonated mono-, di-, and polysaccharides; fatty acids such as palmitic acid, stearic acid, and pleic acid; cholesterol, cholesterol esters, and cholesterol hemisuccinate. Due to their excellent biocompatibility characteristics, phospholipids and combinations of phospholipids and poloxamers are particularly suitable for use in the pharmaceutical embodiments disclosed herein.

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Compatible nonionic detergents comprise: sarbitan esters including sorbitan trialesta (Spen<sup>®</sup> 85), sorbitan sesquioleate, sorbitan monoleate, sorbitan monoleate, polyacyethylene (20) sorbitan monoleate, oleyl polyacyethylene (2) ather, steryl polyacyethylene (4) ather, steryl polyacyethylene (5) ather, steryl polyacyethylene (6) ather, steryl polyacyethylene (6) ather, steryl polyacyethylene and polyacypropylene, including poloxamer 188 (Pluronic<sup>®</sup> F-88), poloxamer 407 (Pluronic<sup>®</sup> F-127), and poloxamar 338. Ionic surfactants such as sodium sudfosuccinate, and fatty acid soaps may also be utilized. In preferred embodiments the microstructures may comprise oleic acid or its alkelical.

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In addition to the dramen-inned surfactants, cationic surfactants or lipids are preferred especially in the case of delivery or RNA or DNA. Exemples of suitable cationic lipids include: DOTMA, N-11-12,3 diolaylaxylpropyll-N.M.N-trimethylammonism chloride; DOTAP, 1,2-dioleyloxy-3-trimethylammoniolpropene; and DOTB, 1,2-dioleyl-3-44-trimethylammoniolprutaneyl-anglycerol. Polycationic emino acids such as solf-visite, and bolyaminine are also contemplated.

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Besides those surfactants enumerated above, it will further be appreciated that a wide range of surfactants may optionally be used in conjunction with the present invention. Moreover, the optimum surfactant or combination thereof for a given application can readily be determined by empirical studies that do not require undue experimentation. Finally, as discussed in more detail below, surfactants comprising the structural matrix may also be useful in the formation of precursor dil-invester emulations (i.e. sprey drying feed stock) used during processing to form the perforated microstructures.

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While such surfectant levels are preferably employed in perforated microstructures, thay may be used to provide stabilized systems comprising relatively anoporous, or substantially solid, periculates. The its, while preferred embodiments will comprise perforated microstructures essociated with high levels of surfectant, acceptable microspheres may be formed using relatively low procesty particulates of the same surfactant concentration (i.e., greater than ebout 20% wolve). In this respect such high surfactant embodiments are specifically contemplated as helm within the scope of the present invention.

In other preferred embodiments, of the invention the structural metrix defining the perforated microstructure optionely compises synthetic on natural polymers or combinations thereof. In this respect useful polymers compises polylecides, polylecides/poratios-polycorides, cyclodestrins, polyacrylates, mathyleilubuse, carboxymathyleilubuse, polylecides, polylecides, polylecides, polylecides, polylecides, polylecides, polylecides, polylecides, polyminity pyrrollicides, polylecides, standards, collegen, galetin, etcl. Examples of polymeric resions that would be useful for the preparation of perforated ink microperticles include: styrene-butadiene, styrene-isoprene, styrene-acrylecides, ethylene-acrylic acid, ethylene-acrylic acid, ethylene-acrylic perforate, styrene-isoprene, styrene-acrylicides, ethylene-acrylic perforated, ethylene-acrylicides, acrylic acid-methyl methacrylata, and vinyl chlarides-iniyal acetals. Those skillad in the art will appreciate that, by selecting the appropriate polymers, the definery efficiency of the perforated microparticles and/or the stability of the dispersions may be tailored to opinitiz the effectiveness of the active or koactive agent.

Besides the aforementioned polymer materials and surfactants, it may be desireble to edd other axcipients to a microsphere formulation to improve particle rigidity, production yield, delivary efficiency and deposition, shelf-life and patient acceptance. Such optional excipients include, but are not limited to: coloning agents, taste masking agents, buffers, hygroscopic agents, analoxidants, and chemical stabilizers. Further, various excipients may be incorporated in, or edded to, the particulate metrix to provide structure and form to the perforated microstructures (6.c. microspheres such as latex particles). In this regard it will be appreciated

that the nightifying components can be removed using a post-production technique such as selective solvent extraction.

Other rigidfying excipients mey include, but are not limited to, carbohydratas including imponseaccharides, disacchanides and polyaacchanides. For exemple, monosaccharides such as detrose (anhydrous and monohydrate), glatectose, manifol, Dramonose, sorticute, sorbose and the like; disacchanides such as tectose, maltose, sucrose, trehalcose, and the like; trisacchanides such as raffinose and the like; and other carbohydrates such as starchas lhydroxyethylstarch), cyclodextins and maltodextrins. Amino acids are also suitable excipients with glycine preferred. Mixtures of carbohydrates and amino acids are further held be within the scope of the present invention. The inclusion of both inorganic le.g. sodium chloride, etc.l. organic selts le.g. sodium citrate, sodium ascorbate, magnesium gluconate, sodium gluconate, tromesthamine hydrochloride, etc.l and buffers is also contemplated. The inclusion of salts and organic solids such as ammonium carbonate, ammonium acetate, ammonium chloride or camphor are also contemplated.

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Yet other praterned embodiments include perforated microstructures that may comprise, or may be coated with changed species that protong residence time at the point of contact or enhance penetration through mucosae. For exempla, anionic changes are known to favor mucosdesion while acclaract readings may be used to essociate that formed microparticulate with negatively changed bioective agents such as genetic material. The charges may be imparted through the association or incorporation of polyanienic or pelycationic materials such as polyacrylic acids, polyvoine, polybectic acid and chitosan.

In addition to, or instead of, the components discussed above, the perforated microstructures will preferably comprise at least one active or bioactive agent. As used herein, the term "active agent" simply refers to a substance that enables the perforated microstructures to perform the desired function. Further, that term "active agent" simply refers to a substance that enables the perforated microstructures to perform the desired function. Further, that term "active agent" shall be held inclusive of the term "bioactive agent" is shall be held to comprise any substance that is used in connection with the diagnosis or treatment of a disease, condition or physiological abnormality in a patient. Perticularly preferred bioactive agents for use in accordance with the invention include anti-allergics, peptides and proteins, pulmonary lung surfactants, bronchodilators and anti-inflammatory steroids for use in the treatment of respiratory disorders such as asthma by inhalation therapy. Preferred active agents for use in accordance with the present invention include pigments, dyes, inks, paints, detargents, food severeterers, agines, adsorbants, absorbents, catalysts, nucleating agents, thicknaring agents, polymars, resins, insulators, fillers, fertilizers, phythohommones, insect pellents, antifouling agents, petricides, fungicides, disinfectants, arefumes, deoderants, and combinations of thereof.

It will be appreciated that the perforated microstructures of the present invention may exclusively comptise one or more active or bloactive agents (a. 100% w/w). However, in selected embodiments the perforated microstructures may incorporate much less bisactive agent depending on the activity thereof. Accordingly, for highly active materials the perforated microstructures may incorporate as little as 0.001% by weight although a

concentration of greater than about 0.1% w/w is prefarred. Other embedments of the immention may comprise greater than about 5%, 10%, 15%, 20%, 25%, 30% or even 40% w/w active or bisactive agent. Still more prefarably the perforated microstructures may compose greater than about 50%, 60%, 70%, 50%, 60% or even 50% w/w active or bisactive agent. The precise amount of active or bisactive agent incorporated in the perforated microstructures of the present invention is dependent upon the agent of choice, the required doze, and the form that agent actually used for incorporation. Those skilled in the art will appreciate that such determinations may be made by using well-known plummacological techniques in combination with the teachings of the present invention.

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With regard to pharmaceutical preparations, any bioactive agent that may be formulated in the disclosed perforated microstructures is expressly held to be within the scope of the present invention. In particularly preferred embodiments, the selected bioactive agent may be edministered in the form of an eerosolized medicements. Accordingly, perticularly compatible bioactive agents comprise any drug that may be formulated as a flouvable dry powder or which is relatively insolable in selected dispersion media. In addition, it is preferred that the formulated agents are subject to pulmonary or asad uprake in physiologically effective amounts. Compatible bioactive agents compise hydrophilic and fipophilic respiratory agents, pulmonary surfactents, bronchodilators, entibiotics, enthérids, enti-inflammatories, staroids, entibisteris, leuketriene inhibitors or entegorists, enticholinargics, entimedoptatics, enembed including DNA and RNA, viral vectors, immunective agents, imaging agents, vaccines, immunesuppressive agents, peptides, protatis and combinations themed. Perticularly preferred bioactive agents for inhalation therety comprise mast call inhibitors (enti-altergics), bronchodilators, and enti-inflammatory steroids such as, for example, cromoplycate (e.g. the sodium sett), and albuterol (e.g. the sulfata salt).

Mora spacifically, exemplary medicaments or bisoctive agents may be selected from, for example, analgasics, e.g. codeine, dilydcomorphine, ergotamine, fentanyl, or morphine, anginal preparations, e.g. diliaemm mast cell inhibitors, e.g. comodyn sodium; antiinfactives, e.g. cephalespoints, macroidest, quiante, pericilias, streptomycin, sulphonamides, tetracyclines and pentamidine; antibistamines, e.g. methapyrilene; entiralitiamentorias, e.g., fluticasoes propionate, betomethasone dipropionate, flunisolide, budesonide, tripedane, cortisone, prednisone, prednisione, prednisione, dezamenthasone, betamethasone, or triamdinolone acetonide; antitussives, e.g. noscapine; bronchodiletors, e.g. ephadrine, advandime, fenoturol, formoterol, isoprenaline, metaproteranol, salbutanol, albutarol, salmeterol, terbutaline; duretics, e.g. emiloride; anticholinergics, e.g. pipatopium, atropine, or oxitropium; lung surfactants e.g. Surfaxia, Exosurf, Survente; ranthines, e.g. aminophylline, theophylline, caffeine; therapeutic proteins and peptides, e.g. DNAse, insufin, glucagoa, LHRH, nafaratin, gossrelin, leurproide, interferon, rhu II-1 receptor, macrophage activation factors such as ylymphokines and murramyl dispetides, opicid peptides and neuropeptides such as enkaphalins, endophins, reain inhibitors, chelacystokinias, DNAse, growth hormones, leutwotinen inhibitors and the like. In addition, bioactive agents that comprise an RNA or ONA sequence, perticularly those useful for gene therapoy, genetic vaccination, genetic telanzation or antisense applications, may be incorporated in the disclosed disparsions as

described herein. Representative DNA plasmids include, but are not limited to pCMVB levelable from Genzyme Corp, Framington, MAI and pCMV-B-gal (a CMV promotor linked to the E. cdi Lec Z gene, which codes for the enzyme B-pelactosidasel.

In any event, the selected active or bisective agent(s) may be associated with, or incorporated in, the perforated microstructures in any form that provides the desired efficacy and is compatible with the choses a production techniques. As used herein, the terms "essociate" or "associating" mean that the structural matrix or perforated microstructure may comprise, incorporate, adsord, absorb, be coated with or be formed by the active or bisociative agent. Where appropriate, the actives may be used in the form of salts [e.g. alkali metal or amine salts or as acid addition salts) or as esters or as solvetes libyriated. In this regard the form of the active or bisoactive agents may be selected to optimize the activity and/or stability of the actives and/or to minimize the solubility of the agent in the suspension medium end/or to minimize perficil appreciation.

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It will further be appreciated that the perferend microstructures according to the invention may, if desired, contain a combination of two or more active ingredents. The agents may be provided in combination in a single species of perforated microstructure or individually in separate species of perforated microstructure or individually in separate species of perforated microstructures. For example, two or more active or bioactive agents may be incorporated in a single feed stock preparation and spray dried to provide a single microstructure species comprising a plurelity of active agents. Conversely, the individual actives could be added to separate stocks and spray dried separately to provide a plurality of microstructure species with different compositions. These individual species could be added to the suspansion medium or dry powder dispensing compartment in any desired proportion and placed in the aerosol delivery system as described below. Further, as alluded to above, the perforated in the international control of the combined with one or more convenience (a.e., a micronized drougl active or bloactive agents to provide the desired dispersion stability or powder dispersibility.

Based on the foregoing, it will be appreciated by those skilled in the ert that a wide variety of active or bioactive agents may be incorporated in the disclosed performed microstructures. Accordingly, the list of preferred active agents above is exemplary only and not intended to be limiting. It will also be appreciated by those skilled in the art that the proper amount of bioactive agent and the timing of the desages may be determined for the formulations in accordance with a leady existing information and without undoe experimentation.

As seen from the passages above, various components may be associated with, or incorporated in the perforated microstructures of the present invention. Similarly, soweral techniques may be used to provide particulates having the desired morphology (e.g. a perforated or hollowyborous configuration), dispersibility and density. Among other methods, perforated microstructures competible with the instant invention may be formed by techniques including spray drying, vacuum drying, solvent extraction, enulainfaction or lyosphilization, and combinations thereof. It will further be appreciated that the basic concepts of many of these techniques are well known in the prior and any would not, in view of the teachings herein, require undue experimentation to adapt them so as to provide the distinct perforated microstructures.

While several procedures are generally competible with the present invention, particularly preferred embodiments typically comprise perferated microstructures formed by spray drying. As is well known, spray drying is a one-step process that converts a liquid feed to a dried particulars form. With respect to pharmaceutical applications, it will be appreciated that spray drying has been used to provide powdered material for various administrature routes including inhalation. See, for example, M. Sacchetti and M.M. Van Oort in: Inhalation Areatols: Physical and Biological Basis for Therapy, A.J. Hickey, ed. Marcel Dekkar, New York, 1996, which is incorporated Papin by reference.

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In general, spray drying consists of bringing together a highly dispersed liquid, and a sufficient volume of hot air to produce evaporation and drying of the liquid droplets. The preparation to be spray dried or feed (or feed stock) can be any solution, course suspension, stury, calloided dispersion, or paste that may be atomized using the selected spray drying apparatus. In preferred embodiments the feed stock will comprise a colloidel system such as an emulsion, reverse emulsion, microemulsion, multiple emulsion, particulate dispersion, or stury. Typically the feed is sprayed into a current of warm filtered air that evaporates the solvent and conveys the dried product to a collector. The spent air is then exhausted with the solvent. Those skilled in that art will appreciate that several different types of apparatus may be used to provide the desired product. For example, commercial spray dryers manufactured by Buchi Ltd. or Niro Corp. will effectively orroduce a pricles of desired stax.

It will further be appreciated that these spray dryers, and specifically their atomizers, may be modified or customized for specialized applications, i.e. the simultaneous spraying of two solutions using a double nozzle technique. More specifically, a water-in-oil emulsion can be atomized from one nozzle and a solution containing an anti-adherent such as manifol can be co-atomized from a second nozzle. In other cases it may be desirable to push the feed solution though a custom designed nozzle using a high pressure liquid chromatography (HPLC) pump. Provided that microstructures comprising the correct morphology and/or composition are produced the choice of apparettus is not critical and would be apparent to the skilled artisan in view of the teachings herein.

While the resulting spray-dried powdered particles typically are approximately spherical in shape, nearly uniform in size and frequently are hollow, then may be some degree of irregularity in shape depending upon the incorporated medicament end the spray drying conditions. In many instances dispersion stability and dispersibility of the perforated microstructures appears to be improved if an inflating agent (or blowing agent) is used in their production. Particularly preferred embodiments may comprise an emulsion with the inflating agent as the disperse or continuous phase. The inflating agent is preferably dispersed with a surfactant solution, using, for instance, a commercially available microfluidizer at a pressure of about 5000 to 15,000 psi. This process forms an emulsion, preferably stabilized by an incorporated currectent, typically comprising submicrom rights of water immiscible blowing agent dispersed in an equeues continuous phase. The formation of such emulsions using this and other techniques are common and well known to those in this art. The blowing agent is

preferably a fluorinatad compound (e.g. parfluorahexane, perfluorocyty bromide, perfluorahezalin, perfluorochtyf ethanel which vaponizes during the spraydrying process, leaving behand generally hollow, porous aerodynamically light microsphares. As will be discussed in more detail bollow, other suitabla liquid blowing agents include nonfluorinated oils, chloroform, Freons, ethyl acetate, alcohols and hydrocarbons. Nitrogen and carbon dioxide gassa are also contemplated as a suitable blowing agent.

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Begides the aforementioned compounds, inorganic and organic substances which can be removed under reduced pressure by sublimation in a post-production step are also competible with the instant invention. These sublimating compounds can be dissolved or dispersed as micronized crystals in the spray drying feed solution and include ammonium carbonate and camptor. Other composition compatible with the present invention comprise rigidifying solid structures which can be dispersed in the feed solution on prepared in-situ. These structures are then extracted after the initial particle generation using a post-production solvent extraction step. For example, latex particles can be dispersed and subsequently dried with other wall forming compounds. followed by extraction with a suitable solvent.

Although the perforated microstructures are preferably formed using a blowing agent as described above, it will be appreciated that, in some instances, no additional blowing agent is required and an aquaous dispersion of the medicament and/or excipients and surfactantls! are spray fried directly. In such cases, the formulation may be amenable to process conditions (a.g., elavated temperatures) that may lead to the formetion of hollow, relatively porous microparticles. Moreover, the medicament may possess special physiochamical proparties (e.g., high crystallinity, elevated melting temperature, surface activity, atc.) that makes it particularly suitable for use in such techniques.

When a blowing agent is employed, the degree of porosity and dispersibility of the perforated microstructure appears to depend, at least in part, on the nature of the blowing agent, its concentration in the feed stock (e.g., as an emulsion), and this spray drying conditions. With respect to controlling porosity and, in suspensions, dispersibility it has surprisingly been found that the use of compounds, heretofore unappreciated as blowing agents, may provide perforated microstructures having particularly desirable characteristics. More particularly, in this novel and unexpected aspect of the present invention it has been found that the use of fluorinated compounds having relatively high boiling points (i.e. greater than about 40°C) may be used to produce particulates that are particularly porous. Such perforated microstructures are especially suitable for inhalation therapies. In this regard it is possible to use fluorinated or partially fluorinated blowing agents having boiling points of greater than about 40°C, 50°C, 60°C, 70°C, 80°C, 80°C, 80°C are away 50°C. Particularly preferred blowing agents having boiling points greater than the boiling point of water, i.e. greater than 100°C (e.g. perflubron, perfluorodacalini). In addition blowing agents with relatively low water solubility (< 10° Mi) are preferred since they enable tha production of stable emulsion dispersions with mean weighted particle diameters less than 0.3 un.

As previously described, these blowing egents will preferably be incorporated in an emulsified feed stock prior to spray drying. For the purposes of the present invention this feed stock will also preferably comprise one or more active or bioactive agants, one or more surfactants of one or more excipients. Of course, combinations of the aforementioned components are also within the scope of the invention. While high boiling (> 100°C) flourinated blowing agents comprise one preferred aspect of the present invention, it will be appreciated that nonfluorinated blowing agents with similar boiling points (> 100°C) may be used to provide perforated microstructures. Exemplary nonfluorinated blowing agents suitable for use in the present invention comprise the formula:

#### R1-X-R2 or R1-X

wherein: R<sup>1</sup> or R<sup>2</sup> is hydrogen, alkyl, alkenyl, alkynl, eromatic, cyclic or combinations thereof, X is any group containing carbon, sulfur, nitrogen, halogens, phosphorus, exvgen and combinations thereof.

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While not limiting the invention in any way it is hypothesized that, as the aqueous feed component evaporates during sprey drying it leaves a thin crust at the surface of the particle. The resulting particle wall or crust formed during the iridial moments of sprey drying appears to trap any high boiling blowing agents as hundrads of amulsion droplets (ca. 200. 300 nml. As the drying process continues, the pressure inside the particulate increases thereby vaporizing at least part of the incorporated blowing agent and forcing it through the relatively thin crust. This venting or outgassing appearantly leads to the formation of pores or other defects in the microstructure. At the same time remaining particulate components [possibly including same blowing agentl migrate from the interior to the surface as the particle solidifies. This migration apparently slows during the drying process as a result of increased resistance to mass transfer caused by an increased internal viscosity. Once the migration ceases the particle solidifies, leaving vaids, poss, defects, hollows, spaces, intertraids spaces, apartures, perforations or holes. The number of pores or detects, their size, and the resulting well thickness is largely dependent on the formulation and/or the nature of the selected blowing agent (e.g. boiling point), its cencentration in the emulsion, total oddic cencentration, and the sprey-drying conditions. It can be greatly appreciated that this type of particle morphology in part contributes to the improved powed refspersibility, suspension stability and exerdynamics.

It has been surprisingly found that substantial amounts of these relatively high bailing blowing agents may be retained in the resulting spray dried product. That is, spray dried perforated microstructures as described herein may comprise as much as 1%, 3%, 5%, 10%, 20%, 30% or even 40% wolve of the blowing agent. In such cases, higher production yields were obtained as a result an increased particle density caused by residual blowing agent. It will be appreciated by those skilled in the art that retained fluorinaed bipowing agent may after the surface characteristics of the perforated microstructures, thereby minimizing particle agengagation during processing and further increasing dispersion stability. Residual fluorinated blowing agent in the particle may also reduce the cohesive forces between particles by providing a barrier or by attenuating the attractive forces produced during manufacturing (e.g., electrostatics). This reduction in cohesive forces

may be particularly advantageous when using the disclosed microstructures in conjunction with dry powder inhalers.

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Furthermore, the amount of residual blowing agent can be attenuated through the process conditions (such as outlet temperature), blowing agent concentration, or boiling point. If the cutet temperature is at or above the boiling point, the blowing agent escapes the particle and the production yield decreases. Preferred outlet temperature will generally be operated at 20, 30, 40, 50, 50, 70, 80, 90 or even 100°C lass than the blowing agent boiling point. More preferably the temperature differential between the outlat temperature and the boiling point will range from 50 to 150°C. It will be appreciated by those skilled in the art that particle porosity, production yield, electrostatics and dispessibility can be optimized by first identifying the range of process conditions (e.g., outlet temperature) that are suitable for the selected active agents and/or excipients. The preferred blowing agent can be then chosen using the maximum outlet temperature such that the temperature differential with be at least 20 and up to 150°C. In some cases, the temperature differential can be outside this range such as, for example, when producing the particulates under supertricial conditions or using lyephilization techniques. These skilled in the art will further appreciate that the preferred concentration of blowing agent can be determined experimentally without under experimentation using techniques similar to those described in the Examples harein.

While residual blowing agent may be advantageous in selected embodiments it may be desirable to substantially remove any blowing agent from the sprey dried product. In this respect, the residual blowing agent can easily be removed with a post-production evaporation step in a vacuum oven. Moreover, such post production techniques may be used to provide perforations in the particulates. For exemple, pones may be formed by spray drying a bloactive agent and an excipient that can be removed from the formed perticulates under a vacuum.

In any event, typical concentrations of blowing agent in the feed stock are between 2% and 50% viv. and more preferably between about 10% to 45% viv. In other embodiments blowing agent concentrations will preferably be greater than about 5%, 10%, 15%, 20%, 25% or aven 30% viv. Yat other feed stock emulsions may comprise 35%, 40%, 45% or even 50% viv of the selected high boiling point companied.

In preferred embodiments, another method of identifying the concentration of blowing agent used in the faed is to provice it as a ratio of the concentration of the blowing agent to that of the stabilizing surfactant (a.g. phosphatidylcholine or PCI) in the precursor or feed emulsion. For fluorocarbon blowing agents (e.g. perfluoroccyt) bromide), and for the purposes of explenation, this ratio has been termed the PFCIPC ratio. More generally, it will be appreciated that compatible blowing agents and/or surfactants may be substituted for the exemplary compounds without falling outside of the scope of the present invantion. In any event, the typical PFCIPC ratio viull range from about 1 to about 60 and more preferably from about 10 to about 50. For preferred embodiments the ratio will generally be greater than about 5, 10, 20, 25, 30, 40 or

even 50. In this respect, Fig. 1 shows a series of pictures taken of perforated microstructures formed of phosphatidylcholine (PC) using various amounts of perfluorocetyl branide (PFC), or relaively high boiling point fluorocethon as the blowing agent. The PFCIPC ratios are provided under each subset of pictures, i.e. from 1A to 1F. Formation and imaging conditions are discussed in greater detail in Examples I and II below. With regard to the micrographs, the column on the laft shows the intact microstructures while the column on the right illustrates cross-sections of frectured microstructures from the same preparations.

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As may easily be seen in the Fig. 1, the use of higher PFC/PC raises provides structures of a more hollow and porous nature. More particularly, those methods employing a PFC/PC raise of grawer than about 4.8 tended to provide structures that are particularly competible with the dry power formulations and dispersions disclosed herein. Similarly, Fig. 3, a micrograph which will be discussed in more detail in Example XII below, illustrates a proferably perous morphology obtained by using higher boiling point blowing agents (in this case perferoedecish).

While relatively high boiling point blowing agents comprise one preferred aspect of the instant invention, it will be appreciated that more conventional and unconventional blowing or inflating agents may also be used to provide compatible perforated microstructures. The blowing agent comprises any volatile substance, which can be incorporated into the feed solution for the purpose of producing a perforated feam-fixe structure in the resulting dry microspheres. The blowing agent may be removed during the initial drying process or during a past production step such as vacuum drying or solvent extraction. Suitable initial drying process or during a

- Dissolved low-boiling (below 100 C) agents miscible with aqueous solutions, such as methylana chloride, acatone, ethyl acetate, and alcohols used to saturate the solution.
- A gas, such as CO<sub>2</sub> or N<sub>2</sub> or liquid such as Freens, CFCs, HFAs, PFCs, HFCs, HFBs, fluoroalkanes, and hydrocarbons used at elevated pressure.
- Emulsions of immiscible low-boding (below 100 C) liquids suitable for use with the present invention are generally of the formula:

R'-X-R2 or R'-X

- wherein: R<sup>1</sup> or R<sup>2</sup> is hydrogen, alkyl, alkenyl, alkynl, aromatic, cyclic or combinations thereof, X is any groups containing carbon, sulfur, nitrogen, halogens, phosphorus, oxygen and combinations thereof. Such liquids include: Freens, CFCs, HFAs, PFCs, HFDs, fluoroalkenss, and hydrocarbons.
- Dissolved or dispersed salts or organic substances which can be removed under reduced pressure by sublimation in a post-production step, such as anymonium salts, camphor, atc.
- Dispersed solids which can be extracted after the initial particle generation using a post-production solvent extraction step, such particles include latex, etc.

With respect to these lower boding point inflating agents, they are typically added to the feed stock in quantities of about 1% to 40% v/v of the surfactent solution. Approximately 15% v/v inflating agent has been found to anoduce a soney died powder that may be used to form the stabilized dispersions of the present invention.

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Regardless of which blowing agent is ultimately selected, it has been found that compatible parforated microstructures may be produced particularly afficiently using a Büchi min's spray drier immodel B- 191, Switzerlandi. As will be appreciated by those skilled in the art, the inlet temperature and the outlet temperature of the spray drier are not critical but will be of such a leval to provide the desired particle size and to result in a product that has the desired activity of the medicament. In this regard, the inlet and outlet temperatures are edjusted depending on the melting characteristics of the formulation components and the composition of the feed stock. The inlet temperature may thus be between 60°C and 170°C, with the outlet temperatures of about 40°C to 120°C depending on the composition of the feed and the desired particulate characteristics. Preferably these temperatures will be from 90°C to 120°C for the inlet and from 80°C to 90°C for the outlet. The flow rate which is used in the spray drying equipment will generally be about 3 ml per minute to about 15 ml per minute. The atomizer air flow rate will vary between values of 25 liters per minute to about 15 ml per minute. Commercially available spray dryers are well known to those in the art, and suitable settings for any particular dispersion can be readily determined through standard empirical testing, with due reference to the exemples that follow. Of course, the conditions may be edjusted so as to preserve biological activity in leaveer melecules such as proteins or peptides.

Though the perforated microstructures are preferably formed using fluorinated blowing agents in the form of an emulsion, it will be appreciated that nonfluorinated dils may be used to increase the loading capacity of active or bioactive agents without compromising the microstructure. In this case, selection of the nonfluorinated oil is based upon the solubility of the active or bioactive agent, water solubility, boiling point, and flash point. The active or bioactive agent will be dissolved in the oil and subsequently emulsified in the feed solution. Preferably the oil will have substantial solubilization capacity with respect to the selected agent, low water solubility (< 10°M), boiling point greater than water and a flash point greater than the drying outlet temperature. The addition of surfactants, and co-solvents to the nonfluorinated oil to increase the subublication capacity within the scope of the present invention.

In particularly preferred embodiments nonfluorinated oils may be used to solubilize agents or bioactive agents that have limited solubility in equeous compositions. The use of nonfluorinated oils is of particular use for increasing the loading capacity of steroids such as beclomethasone dipropriants of triamoniculone acatonide. Preferably the oil or oil mixture for solubilizing these clathrate forming steroids will have a refractiva index between 1.36 and 1.41 (e.g. athyl butyrata, butyl carbonate, dibutyl ether). In addition, process conditions, such as temperature and pressure, may be adjusted in order to boost solubility of the selected agent. It will be appreciated that selection of an appropriate oil or oil mixtures and processing

conditions to maximize the loading capacity of an agent are well within the purview of a skilled artisan in view of the teachings harein and may be accomplished without undue experimentation.

Particularly preferred embodiments of the present invention comprise spray dying preparations comprising a surfectant such as a phospholipid and at least one active or bioactive agent. In other embodiments the spray drying preparation may further comprise an excipient comprising a hydrophilic moistly such as, for example, a carbohydrate i.e. glucose, lactose, or starch in addition to any selected surfactant. In this regard various starches and derivatized starches suitable for use in the present invention. Other optional components may induce conventional viscosity modifiers, buffers such as phosphate buffers or other conventional biocomposible buffers or phi adjusting agents such as ecids or bases, and compotic spents (to provide isotonicity, hyperomalarity, or hyposmolarity). Examples of stitable selts include sodium phosphate (both monobastic and dibasic), sodium chloride. resicium hospathus, calcium chloride and other drivisolocically acceptable selts.

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Whatever components are selected, the first step in particulate production typically comprises feed stock preparation. Perferably the selected drug is dissalved in water to produce a concentrated solution. The drug may also be dispersed directly in the emulsion, particularly in the case of water insoluble spents. Alternatively, the drug may be incorporated in the form of a solid particulate dispersion. The concentration of the active or bisoctive agent used is dependent on the amount of particulate dispersion. The concentration of the active or bisoctive agent used is dependent on the amount of particulate dispersion. The concentration of the active or bisoctive agent used is dependent on the amount of particulate dispersion. All provides and the performance of the delivery device amployed (e.g., the fine particula dose for a MDI or DPI). As needed, cosurfactants such as polexamer 188 or span 80 may be dispersed into this annex solution. Additionally, excisients such as sources and storches can also be added.

In selected embodiments an oil-in-water emulsion is then formed in a separate vessel. The oil amployed is preferably a fluorocarbon (e.g., perfluorocarbon termine), perfluorodecalini which is emulsified using a surfactant such as a long chain saturated phospholipid. For example, one gram of phospholipid may be homogenized in 150 g bot distilled water (e.g., 80°C) using a suitable high shear mechanical mixer (e.g., Ulturar model T-25 mixer) at 8000 rpm for 2 to 5 minutes. Typically 5 to 25 g of fluorocarbon is adoptive to the dispersed surfactant solution while mixing. The resulting perfluorocarbon in water emulsion is then processed using a high pressure homogenizer to reduce the partial size. Typically the emulsion is processed at 12,000 to 18,000 psi, 5 discreta passes and kept at 50 to 80°C.

The active or bioactive agent solution and perfluencearbon emulsion are then combined and fef into the spray dryer. Typically the two preparations will be miscible as the emulsion will preferrebly comprise acqueous continuous phase. While the bioactive agent is solubilized separately for the purposes of the instant discussion it will be appreciated that, in other embodiments, the active or bioactive agent may be solubilized (or dispersed) directly in the emulsion. In such cases, the active or bioactive emulsion is simply spray dried without combining a separate drug preparation.

In any event, operating conditions such as inlet and outlet temperature, feed rete, atomization pressure, flow rate of the drying air, and nozzle configuration can be adjusted in accordance with the

manufacturer's guidelines in order to produce the required particle size, and production yield of the resulting dry microstructures. Exemplary settings are as follows: an air inlet temperature between 60°C and 170°C; an air outlet between 40°C to 120°C; a feed rate between 3 ml to about 15 ml per minute; and an aspiration air flow of 300 L/min. and an atomization air flow rate between 25 to 50 L/min. The selection of appropriate apparatus and processing conditions are well within the puriew of a skilled artisan in view of that tacchings herein and may be accomplished without undue experimentation. In any avent, the use of these and substantially equivalent methods provide for the formation of hollow perous aerodynamically light microspheres with particle diameters appropriate for aerosol deposition into the lang, microstructures that are both hollow and porous, almost honeycombad or foam-fike in appearance. In especially preferred anhadiments the perforated microstructures commons hollow, no your survay dried microsobstres.

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Along with sprey drying, perferented microstructures useful in the present invention may be formed by hysphilization. Those skilled in the art will epprecise that tyophilization is a freeze drying process in which water is sublimed from the compestion after it is frozen. The pericular edventage associated with the lyophilization process is that biologicals and pharmaceuticals that are relatively unstable in an equeous solution can be dried without elevated temperatures (thereby eliminating the adversa thermal affacts), and than storad in a dry state where there are few stability problems. With respect to the instant invantion such techniques are perticularly compatible with the incorporation of peptides, protains, genetic material and other natural and synthetic macromolecules in particulates or perforated microstructures without compromising physiological activity. Mathods for providing lyophilized particulates are known to those of skill in the art and it would clearly not require undue exparimentation to provide dispersion compatible microstructures in accordance with the teachings herein. The lyophilized cabe containing a fine foam-like structure can be micronized using techniques known in the art to provide 3 to 10µm sized particles. Accordingly, to the extent that lyophilization processes may be used to provide microstructures having the desired poresity and size they instant invention.

Besides the aforementioned techniques, the perforated microstructures or particlas of the present invention may also be formed using a method where a feed solution (either emulsion or aqueous) containing wall forming agents is rapidly added to a meservoir of heated oil (e.g. perflubron or other high boiling FCal under reduced pressure. The water and valutie solvants of the feed solution rapidly boils and are evaporated. This process provides a perforated structure from the wall forming agents similar to purified rice or popcorn. Preferably the wall forming agents among perticles can then separated from the heated oil using a filtering technique and subsequently dried under vacuum.

Additionally, the perforeted microstructures of the present invention may also be formed using a double emulsion method. In the double emulsion method the medicament is first dispersed in a polymer dissolved in an organic solvent (e.g. methylene chloride) by sociacion or homogenization. This primary

emulsion is then stabilized by forming a multiple emulsion in a continuous aqueous phase containing an emulsifier such as polyvinylelochal. Evaporation or extraction using conventional techniques and apparatus then removes the organic solvent. The resulting microspheres are washed, filtered and dried prior to combining them with an appropriate suspension medium in accordance with the present invention

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Whatever production method is utimately selected for production of the perforated microstructures, the resulting powders have a number of advantageous properties that make them pericularly compatible for use in devices for inhalation tharapies. In particular, the physical characteristics of the perforated microstructures make them extremely effective for use in dry powder inhales and in the formation of stabilized dispersions that may be used in conjunction with metared dose inhalars, nebulizers and liquid dose instillation. As such, the perforated microstructures provide for the effective pulmonary administration of livers in a postate.

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In order to maximize dispersibility, dispersion stability and optimize distribution upon administration, the mean geometric particle size of the perforated microstructures is preferably about 0.5-50 m. more preferably 1-30 m. It will be appreciated that large particles (i.e. preseter than 50 mi may not be postered in applications where a valve or small oiffice is employed, since large particles tend to aggregate or separate from a suspension which could potentially clog the device. In especially preferred embodiments the mean geometric particle size for diametar) of the perforated microstructures is lass than 20 m or less than 10 m. More prisarely the mean geometric diameter is less than about 7 m or 5 m, and even more preferably less than about 2.5 m. Other preferred ambodiments will comprise proparations wherein the mean geometric diameter of the perforated microstructures is between about 1 m and 5 m. In especially preferred embodiments the perforated microstructures will comprise a powder of dry, hollow, porous microsphesical shells of approximately 1 to 10 m or 1 to 5 m in dameter, with shell thicknesses of approximately 0.1 m to approximately 0.5 m. It is a particular advantage of the present invention that delivery characteristics of the selected particle size.

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As alluded to throughout the instant specification the porceity of the microstructures may play a significant part is establishing dispersibility (e.g., in DPIs) or dispersion stability (e.g., for MDIs or nebulizers). In this respect, the mean porceity of the perforated microstructures may be determined through abecton microscopy coupled with modern imaging techniques. More specifically, electron microgrephs of representative samples of the perforated microstructures may be obtained and digitally analyzed to quantify the perceity of the preparation. Such methodology is well known in the ert and may be understated without undoe experimentation.

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For the purposes of the present invention, the mean porceity file, the percentage of the particle surface area that is open to the interior and/or a central void of the perforated microstructures may range from approximately 0.5% to approximately 80%. In more preferred embodiments, the mean proxify will range from approximately 2% to approximately 40%. Based on selected production parameters, the mean pomisiry may be presented than approximately, 2% 5%, 10%, 15%, 20%, 25% or 30% of the microstructure surface eres. In other

embodiments, the mean parasity of the microstructures may be greater than about 40%, 50%, 60%, 70% or even 80%. As to the powes themselves, they typically range in size from about 5 mm to about 400 nm with mean pore sizes preferably in the range of from about 20 nm to about 200 am. In particularly preferred embodiments the mean pore size will be in the range of from about 50 rm to about 100 nm. As may be seen in Figs. 1Alt to 1F2 and discussed in more detail below, it is a significant edventage of the present invention that the pore size and porocity may be closely controlled by careful selection of the incorporated components and production parameters.

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In this regard, the particle morphology end/or hollow design of the perforeted microstructures also plays an import ent role on the dispersibility or collectiveness of the dry provider formulations disclosed bravin. That is, it has been surprisingly discovered that the inherent cohesive character of fine providers can be overcome by lovening the world review of Weaks, electrostatic attractive and fliquid bidging forces that typically exist between dry particles. More specifically, in concordance with the teachings herein, improved provider dispersibility may be provided by engineeing the particle morphology and density, as well as control of humidity and charge. To that end, the perforated microstructures of the present invention comprise pores, wide, hollows, defects or other intersitial spaces which reduce the surface contact area between particles thereby minimizing interpractic forces. In addition, the use of surfacetants such as phospholipids and fluoriested blowing agents in accordance with the teachings herein may contribute to improvements in the flow properties of the powders by tempering the charge and strength of the electrostatic forces as well as misture content.

Most fire poviders (e.g. < 5 /ml exhibit poor dispersibility which can be problematic when attempting to deliver, sensolize endlor package the providers. In this respect the major forces which control parties interactions can typically be divided into long and short range forces. Long range forces include gravitational extractive forces and electrostatics, where the interaction varies as a square of the separation distance or particle diameter. Important short range forces for dry poviders include van der Waals interactions, hydrogen bonding and fiquid bindess. The latter two short range forces differ from the others in that they occur where there is already contact between particles. It is a major advantage of the present invention that these attractive forces may be substantially attenuated or reduced through the use of perforted microstructures as described herein.

In an effort to overcome these attractive forces, typical prior at dry powder formulations for DPIs comprise microrized drug particles that are deposited on large cereirs particles (e.g., 20 to 99 Juml) such as lactose or appliamented units of pure drug particles or appliamention of fine lactose particles with pure drug, since they ere more readily fluidized than next drug particles. In addition, the mass of drug required per actuation is typically less than 100 µg and is thus prohibitively too small to meter. Hence, the larger lactose particles in prior art formulations function as both a carrier particle for secrosofization and a bulking agent for metering. The use of large particles in these formulations are employed since powder dispensibility and secrosofization efficiency improves with increasing increasing particle size as a result of diminished interparticle forces (French, D.L., Edwards, D.A., sand fliven, R.W., J. Aerosol Sci. 27, 786-783, 1998 within is increprometed herein by reference). That is, prior art formulations of the use

large particles or carriers to overcome the principle forces controlling disparsibility such as van der Waals forces, liquid bridging, and electrostetic attractive forces that exists between particles.

These skilled in the art will expressive that the way dee Waels (VIDW) attractive force occurs at short reage and depends, at least in part, on the surface context between the interesting particles. When two dry particles approach each other the VIDW forces increase with an increase in contact area. For two dry particles, the magnitude of the VIDW interestion force. P.... can be calculated using the following assistance:

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$$F_{\text{wdw}}^0 = \frac{\hbar \omega}{8\pi d_0^2} \left[ \frac{r_1 r_2}{r_1 + r_2} \right]$$

where h is Planck's constant, to is the angular frequency, d<sub>0</sub> is the distance at which the adhesional force is at a maximum, and r<sub>0</sub> and r<sub>2</sub> are the redii of the two interacting particles. Accordingly, it will be appreciated that one way to minimize the magnitude and strength of the VDW force for dry powders is to decrease the interparticle area of contact. It is important to note that the magnitude d<sub>0</sub> is a reflection of this area of contact. The minimal area of contact between two opposing bodies will occur if the particles are parfect spheres. In addition, the erea of contact will be further minimized if the particles are highly porous. Accordingly, the performed microstructures of the present immention act to reduce interparticle contact and corresponding VDW ettractive forces. It is important to note that this reduction in VDW forces is largely a result of the unique particle morphology of the powders of the present immention rather than an increase in geometric particle diameter. In this regard, it will be appreciated that particularly preferred embediments of the present immention provide powders having average or small particularly preferred embediments of the present immention provide powders having average or small particular (a.g., per particle distinctor of the present immention provide powders having average or small particular (a.g., per particle such as conventional micronized drugs of the same size will exert greater interparticle forces between them and hence, will obtain too or cowder desponsibility.

Further, as indicated above, the electrostatic force affacting powders occurs when either or both of the particles are electrically charged. This phenomenon will result with either an attraction or repulsion between particles depending on the similarity or dissimilarity of charge. In the simplest case, the electric charges can be described using Coulomb's Law. One way to modulate or decrease the electrostatic forces between particles is if either are both periodes have non-conducting surfaces. Thus, if the perforated microstructure powders comprise excipients, surfactants or active significant shall be a conducted to the period of the period will be uneverly distributed over the surface. As a result, the charge half-life of powders comprising non conducting components will be relatively short since the retention of elevated charges is dictated by the resistivity of the material. Resistive or non-conducting components are materials which will seither function as an efficient electron donor or accordance.

Derjaguin et al. (Murler, V.M., Yushchenko, V.S., end Derjaguin, B.V., J. Callaid Interface Sci. 1890. 77, 115-119), which is incorporated herein by reference, provide a list renking melacular groups for their ebility to accept or donate an elactron. In this report examplery groups may be ranked as follows:

Donor: 
$$\cdot NH_2 > \cdot OH > \cdot OR > \cdot COOR > \cdot CH_3 > \cdot C_8H_5 > \cdot Adogen > \cdot COOH > \cdot CO > \cdot CN Acceptor:$$

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The present invention provides for the reduction of electrostatic effects in the discissed powders though the use of relatively nonconductive metanials. Using the above rankings, preferred nonconductive metenals would include halogranted endoir hydrogeneted components. Metanial such as phesphalipids and fluorinated blowing spents (which may be retained to some extent in the spary dried powders) are preferred since they can provide resistance to particle charging. It will be appreciated that the retention of residual blowing agent (e.g. fluorochemicals) in the particles, even at relatively low levels, may help minimize charging of the perforated microstructures as is typically imparted during spray drying and cyclone separation. Besed on general electrostatic principles and the teachings herrin, one skilled in the ent would be eithe to identify additional materials that serve to reduce the decreastatic forces of the disclosed powders without undue experimentation. Further, if reeded, the electrostatic forces can also be manifolded and minimized using electrification and cheministes.

In addition to the surprising advantages described above, the present invention further provides for the attenuation or reduction of hydrogen and siguid bonding. As known to those skilled in the art, both hydrogen bonding and Siguid bridging can result from moisture that is absorbed by the provider. In general, the provider is a provided in the art beamscautical formulations for inhelation therepies which tend to employ relatively hydrophilic compounds such as licitose. However, in accordance with the teachings herein, adhesion forces due to adsorbed water can be modulated or reduced by increasing the bydrophobicity of the contacting surfaces. One skilled in the ert can approciate that an increase in puricle hydrophobicity can be achieved through excipient selection and/or use a post-production spray drying cosmity technique such as employed using a fluidized bed. Thus, preferred excipients include hydrophobic surfactants such as phospholipids, fatty acid soaps and cholasterd. In view of the teachings herein, it is submitted that a skilled arissan would be able to identify meterials exhibiting similar desirable properties without undue experimentation.

In accordance with the present invention, methods such as engle of repose or shear index can be used to assess the flow properties of dry providers. The engle of repose is defined as the engle farmed when a cane of powder is poured ento a flat surface. Powders having an engle of repose renging from 45° to 20° are preferred end indicate suitable provider flow. More perticularly, powders which passass an engle of repose between 33° and 20° exhibit relatively lows shear forces and are especially useful in pharmaceutical preparations for use in inhalation therapies (e.g. DDT.). The shear index, though more time consuming to measure than engle of repose, is considered

more reliable and easy to determine. Those skilled in the art will appreciate that the experimental procedure outlined by Arridon and Houghton IG.E. Amidon, and M.E. Houghton, Pharm. Manuf., 2, 20, 1985, incorporated herein by reference) can be used estimate the shear index for the purposes of the present invention. As described in S. Kocova and N. Pilpel, J. Pharm. Pharmacol. 8, 33-55, 1973, also incorporated herein by reference, the shear index is estimated from powder parameters such as, yield stress, effective angle of internal friction, tendle strength, and specific cohesion. In the present invention powders having a shear index less than about 0.08 are desirable. More preferably, powders used in the disclosed compositions, methods and systems will have shear indicas less than about 1.1. In particularly preferred embodiments the shear index will be less than about 1.3 or even less than about 1.5. Of course powders having different shear indices may be used provided the result in the effective deposition of the scrive or bioscrive agent at the site of interest.

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It will also be appreciated that the flow properties of powders have been shown correlate well with bulk density measurements. In this regard, conventional prior art thinking (IC. Harwood, J. Pharm, Sci., 60, 161-163, 1671) held that an <u>increase</u> in bulk density correlates with improved flow properties as pradicted by the shelf of the material. Conversely, it has surprisingly been found that, for the perforated microstructures of the present invantion, suprior flow properties were exhibited by powders having relatively how bulk densities. That is, the hollow porous powders of the present invention exhibited superior flow properties over powders substantially devoid of pores. To that and, it has been found that it is possible to provide powders having bulk densities of less than 0.5 glcm<sup>2</sup> that anxibit particularly favorable flow properties. More surprisingly, it has been found that it is possible to provide perforated microstructure powders having bulk densities of less than 0.3 glcm<sup>2</sup> or even less than about 0.1 glcm<sup>2</sup> that exhibit excellent flow properties. The ability to produce low bulk density powders having superior flowability further eccentuates the noval and weapopetal nature of the present invention.

In addition, it will be appreciated that the reduced attractive forces (e.g. van der Wasts, alactrostatic, hydrogen and liquid bonding, etc.) and excellent floweithly provided by the perforated microstructure powders make them perticularly useful in preparations for inhalation therapies (e.g. in inhalation devices such as DHs, MDIs, rebutizers). Along with the superior flowebility, the perforated or porous and/or hellow design of the microstructures also plays an important role in the resulting ceresol properties of the powder when discharged. This phenomenon holds true for perforated microstructures aerosolized as a suspension, as in the case of an MDI or a nebulizer, or delivery of perforated microstructures in dry form as in the case of a OPI. In this respect the perforated articuture and relatively high surface area of the dispersed microparticles anables them to be carried along in the flow of gases during inhalation with greater ease for longer distances than non-perforated nortices of commerciale size.

More pericularly, because of their high porosity, the density of the particles is significantly less than 1.0 g(cm², typically less than 0.5 g(cm², more often on the order of 0.1 g(cm², and as low as 0.01 g/cm². Unlike the geometric particle size, the aerodynamic particle size,  $d_{aer}$ , of the perforated microstructures depends substantially on the particle density,  $\rho: d_{aer} = d_{geo}\rho$ , where  $d_{geo}$  is the geometric diameter.

For a particle density of 0.1 g/cm<sup>2</sup>,  $d_{arr}$ , will be roughly three times smaller than  $d_{gro}$ , leading to increased particle deposition into the psipheral regions of the lung and correspondingly less deposition in the throat. In this regard, the mean serodynamic diameter of the perforated microstructures is preferably less than about 5  $\mu$ m, more preferably less than about 3  $\mu$ m, and, in particularly preferred embodiments, less than about 2  $\mu$ m. Such particle distributions will act to increase the deep lung deposition of the blocative agent whether administered using a OPI, MDI or nebulizer. Further, having a larger geometric diameter than aerodynamic diameter brings the perticles closer to the wall of the alvaolus thus increasing the deposition of small aerodynamic diameter sortices.

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As will be shown subsequently in the Examples, the particle size distribution of the aerosal formulations of the present invention are measurable by conventional techniques such as, for example, esscade impaction or by time of flight analytical methods. In addition, detarmination of the emitted dose from inhalation devices were done according to the proposed U.S. Phermacopeia method (Pharmacopeial Previews, 221986) 30651 which is incorporated herein by reference. These and releted techniques enable the "fine particle frection" of the eerosol, which corresponds to those particulates that are likely to effectively deposited in the lung, to be calculated. As used herein the phrase "fine particle frection" refers to the percentage of the total amount of active medicament delivered per actuation from the mouthpiece of a DPI, MDI or nebulizer onto plates 2-7 of an 8 stage Andersen cascade impactor. Besed on such measurements the formulations of the present invention will preferably have a fine particle fraction of approximately 20% or more by weight of the perforated microstructures (whu), more preferably they will actibit a fine particle fraction of from about 25% to 80% w/w, and even more preferably from about 30 to 70% w/w. In selected medicaments the present invention will preferably comprise a fine particle fraction of greater than about 30%, 40%, 50%, 60%, 70% or 80% by weight.

Further, it has also been found that the formulations of the present invention exhibit relatively low deposition rates, when compared with prior art preparations, on the induction port and onto plates 0 and 1 of the impactor. Deposition on these components is linked with deposition in the throat in humans. More specifically, most commercially available MOIs and DPIs have simulated throat depositions of approximately 40.70% (wive) of the total doss, while the formulations of the present invention typically deposit less than about 20% w/w. Accordingly, preferred embodiments of the present invention heve simulated throat depositions of less than about 40%, 35%, 30%, 25%, 20%, 15% or even 10% w/w. Those skilled in the art wall appreciate that significant decrease in throat deposition provided by the present invention will result in a corresponding decrease in associated local side-effects such as throat initiation and candidasis.

With respect to the advantageous deposition profile provided by the instant invention it is well known that MIDI propellants typically force suspended particles out of the device at a high velocity towards the back of the throat. Since prior art formulations typically contain a significant percentage of large particles and/or sourceates, as much as two-thirds or more of the emitted dose may impact the threat.

Mareavar, the undesirable delivery profile of conventional powder preparations is also exhibited under conditions of low particle velocity, as occurs with DPI devices. In general, this problem is inherent when serosolizing solid, danas, particulates which are subject to aggregation. Yet, as discussed above, the novel and unexpected properties of the stabilized dispersions of the present invention result in surprisingly low throat deposition upon administration from inhalation device such as a DPI, MDI atomizer or nebulizer.

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While not wishing to be bound by any particular theory, it appears that the reduced throat deposition provided by the instant invention results from decreases in particle aggregation and from the hollow and/or parous morphology of the incorporated microstructures. That is, the hollow and porous nature of the dispersed microstructures slows the velocity of particles in the propellent stream for gas stream in the case of DPIsI, just as a hollow/porous whiftle ball decelerates faster than a basebell. Thus, rather than impacting and sticking to the back of the throat, the relatively slow travaling particles are subject to inhelation by the patient. Moreover, the highly porous nature of the particle sellows the propellent within the perforated microstructure to rapidly leave and the particle density to drop before impacting the throat.

Accordingly, a substantially higher percentage of the administrated bioactive agent is deposited in the pulmonary air passages where it may be efficiently obsorbed.

With respect to inhalation therapies, those skilled in the art will appreciate that the perforated nicrostructure powders of the present invention are particularly useful in DPIs. Conventional DPIs, or dry powder inhalers, comprise providered formulations and devices where a predetermined dose of medicanent, aither alone or in a blend with lactose cernier particles, is delivered as a fine mist or excess of dry powder for inhalation. The medicanent is formulated in a wey such that it readly disperses into discrete particles with a size rage between 0.5 to 20 µm. The powder is actuated either by inspiration or by some external dislivery force, such as pressurized air. DPI formulations are typically packaged in single dose units or they employ reservoir systums capable of mataining multiple doses with manual transfer of the dose to the device.

DPIs are generally classified based on the dose delivery system employed. In this respect, the two major types of DPIs comprise unit dose delivery devices and bulk reservoir delivery systems. As used herein, the term "reservoir" shall be used in a general sense and held to encompass both configurations unless otherwise dictated by contextual restraints. In any event, unit dose delivery systems require the dose of powder formulation presented to that device as a single unit. With this system, the formulation is profiled into dose of powder formulation presented to that device as a single unit. With this system, the formulation is profiled into dose question which may be finish packaged or presented in likities strips to prevent moisture impress. Other unit dose packages include hard gelatin cepsules. Most unit dose containers designed for DPIs are filled using a fixed volume technique. As a result, there are physical limitations (hera density) to the minimal dose that can be metered into a unit dose container is in the range of 5 to 15 mg which corresponds to drug loading in the range of 25 to 15 mg which corresponds to drug loading in the range of 25 to 500/cg per dose. Conversely, bulk reservoir delivery systems provide a precise quantity of powder to be matered upon individual delivery for up to approximately 200 doses. Again like the unit dose systems, the

provider is metered using a fixed volume cell or chamber that the powder is filled into. Thus, the density of the powder is a major factor limiting the minimal dose that can be delivered with this device. Currently bulk reservoir type DPIs can meter between 200µg to 20 mg powder per actuation.

DPIs am designed to be manipulated such that they break open the capsule/bister or to load bulk powder during actuation, followed by dispersion from a mouthpiece or actuator due to the patient's inspiration. When the price art formulations are actuated from a DPI device the lactosiding aggregates are secondard and the patient inhales the mist of dry powder. During the inhalation process, the carrier particles encounter shear forces whereby some of the micronized drug particles are separated from the lactose particulate surface. It will be apprecisted that the drug particles are subsequently carried into the lung. The large lactose particles impact the throat and upper airways due to size and inertial force constraints. The efficiency of delivery of the drug particles is dictated by their degree of adhesion with the carrier particles and their send-variant property.

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Deaggregation can be increased through formulation, process and device design improvements. For example fine particle factors (FPU is often mixed with coarse factors carriers, wherein the FPL will occupy high-energy binding sites on the carrier particles. This process provides more passive sites for adhasion to micronized drug particles. This tertiary bland with the drug has been shown to provide statistically significant increases in fine particle fraction. Other strategies include specialized process conditions where drug particles are mixed with FPL to produce agglementated units. In order to further increase particulate deposition, many DPLs are designed to provide deaggregation by passing the desage form over bafflas, or through tortuous channels that disrupts the flow properties.

The addition of FPL, agglomeration with FPL and specialized device design provides an improvement in the deaggregation of formulations, however, the clinically important parameter is the fine particle dose received by the patient. Though improvements in deaggregation can be provided, a major problem still exists with current DPI devices in that there is an increase in respirable dose with an increased disapiratory effort. This is a result of an increased fine particle fraction corresponding to the increased disapgregation of particle applomerates as the airflow increases through the inhaler with increasing inspiratory effort. Consequently design accuracy is compromised, leading to complications when the devices are used to administer highly efficacious drugs to assnitive populations such as children, adelescents and the alderly. Moreover, the dosing inaccuracy associated with conventional preparations could complicate regulatory approval.

In stark contrast, the perforated microstructure powders of the present invention obviete many of the difficulties associated with prior art centire preparations. That is, an improvement in DPI performance may be provided by engineering the periods, size, secrolymanics, morphology and density, as well as control of humidity and change. In this respect the present invention provides formulations wherein the medicement and the incipients or bulking agents are preferably associated with or comprise the perforated microstructures. As set forth above, preferred compositions according to the present invention typically yield powders with bulk densities

less than 0.1 gicm<sup>2</sup> and often less than 0.05 gicm<sup>2</sup>. It will be appreciated that providing powders having bulk denaities an order of a magnitude less than conventional DPI formulations allows for much lower doses of the salected bloactive agent to be filled into a unit dose container or metered via raservoir-based DPIs. The ability to affectively mater small quantities is of particular importance for low dose statrid, long acting branchodilators and naw protain or peptide medicaments proposed for DPI delivery. Moraover, the ability to effectively deliver particulates without associated carrier particles simplifies product formulation, filling and reduces undestinable side affects.

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As discussed above, the hollow porous powders of the present invention exhibit superior flow properties, as measured by the angle of repose or shear index methods described herein, with respect to equivalent powders substantially devoid of pores. That is, superior powder flow, which appears to be a function of bulk density and puricle morphology, is obsarved where the powders have a bulk density less than 0.5 g/cm². Preferably the powders have bulk densities of less than about 0.3 g/cm², 0.1 g/cm² or even less than about 0.05 g/cm². In this regard, it is theorized that the perforated microstructures comprising poras, voids, hollows, defects or other intensities spaces contribute to powder flow properties by reducing the surface contact, allows, defects or other intensities appeared by the properties of the powder flow properties by reducing the surface contact ventures between particles and minimizing interperticle forces. In addition, the use of phospholipids in preferred embodiments and retention of fluerinated blowing agents may also contributs to improvements in the flow properties of the powders by tempering the charge and strength of the alectrostatic forces as well as meisture content.

In addition to the aforementioned advantages, the disclosed powders exhibit its overable serrodynamic properties that make them particularly effective for use in DPIs. More specifically, the parforated structure and relatively high surface area of the microparticles enables them to be carried along in the flow of gassa during inhalation with greater assa and for longer distances than relatively non-perforated particles of comparable size. Because of their high porceity and low deseity, administration of the perforated microstructures with a DPI provides for increased particle deposition into the peripheral regions of the lung and correspondingly less deposition in that throat. Such particle distribution acts to increase the deep lung deposition of the administration and the peripheral regions of the surgent and deposition of the administration and provides for systemic administration. Moreover, in a substantial improvement over prior at DPI preparations the low-density, highly porous powders of the present invantion preferably eliminate the need for carrier particles. Moreover, in a substantial improvement over prior at DPI preparations the low-density, highly porous powders of the present invantion preferably eliminate the need for carrier particles. Since the lung lactose carrier particles will impact the throat and upper sinveys due to their size, the elimination of such particles minimizes throat deposition and any associated "our" affect associated with conventional DPIs.

Along with their use in a dry powder configuration, it will be appreciated that the perforated microstructuras of the present invention may be incorporated in a suspension medium to provide stabilized dispersions. Among other uses, the stabilized dispersions provide for the effective delivery of bioactive agents to the pulmonary air passages of a patient using MDIs, nebulizers or liquid does instillation (LDI techniques).

As with the DPI embediments, Administration of a bisactive agent using an MDI, nebulizer or LDI technique may be indicated for the treatment of mild, moderate or severa, excut or channic symptoms or far prophylectic treatment. Moreover, the bisactive agent may be edministrated to treat local or systemic conditions or disorders. It will be appriciated thet, the precise does edministrated will depend on the ege and condition of the patient, the pericular medicament used and the frequency of edministration, and will uttimately be at the discretion of the attendant physician. When combinations of bisective agents are employed, the dose of each component of the combination will centerally be that employed for each component when used abone.

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These skilled in the ert will appreciate the enhanced stability of the disclosed dispersions or suspansions is largely exhived by lowering the van der Weals attractive forces between the suspanded particles, and by reducing the differences in density between the suspansion medium and the perticles. In eccordance with the teachings herein, the increases in suspansion stability may be imparted by engineering perforated microstructures which are then dispersed in a competible suspansion medium. As discussed above, the perforated microstructures comprise pores, voids, hollows, defects or other interstitial spaces that allow the fluid suspansion medium to fraely permeate or perfuse the perticulate boundary. Perticularly preferred embodiments comprise perforated microstructures that are both hollow and persus, almost honeycombed or foam-list opposerons. In especially preferred embodiments the perforated microstructures commiss hollows. Dorous serve viried microsotheres.

When the perforated microstructures are placed in the suspension medium fi.e. propellant, the suspension medium is able to permeate the perticles, thereby creeing e "homodispersion", wherein both the sontinuous and dispersed phases ere indistinguishable. Since the defined or "virtual" particles fi.e. comprising the volume circumscribed by the microperticulate matrix) are made up almost entirely of the medium in which they are suspended, the forces driving perticle aggregation (floculation) ere minimized. Additionally, the differences in density between the defined particles and the continuous phase are minimized by having the microstructures filled with the medium, thereby effectively slowing particle creaming or sedimentation. As such, the perforated microspheres and stabilized suspensions of the present invention are particularly compatible with many aerosolization techniques, such as MOI and nebulization. Moreover, the stabilized dispersions may be used in flouid dose instillation applications.

Typical prior art suspensions le.g. for MDIsi comprise mostly solid particles and small amounts (< 1% wilv) of surfactant (e.g. lecitin, Span95, olici acid) to increase electrostatic repulsion between particles or polymers to stancially decrease particle interaction. In sharp contrast, the suspensions of the present invention are designed not to increase repulsion between particles. but rather to decrease the structive forces between particles. The principal forces driving flocculation in nonequeous media are van der Weals attractive forces. As discussed above, VDW forces are quantum mechanical in origin, and can be visualized as attractions between fluctuating dipoles (i.e. induced dipole-induced dipole interactional. Dispersion forces are extremely short-range and scale as the sixth power of the distence between atoms.

When two macroscopic bodies approach one another the dispersion attractions between the atoms sums up.
The resulting force is of considerably longer range, and depends on the geometry of the interacting bodies.

Mere specifically, for two spherical particles, the magnitude of the VDW potential,  $V_A$ , can be approximated by:  $v_A = \frac{-A_{off}}{6A_{off}} \frac{R_i R_i}{(R_i + R_i)}$ , where  $A_{off}$  is the effective Hamaker constant which accounts for the nature of the particles and the medium,  $H_0$  is the distance between particles, and  $R_1$  and  $R_2$  are the radii of spherical particles 1 and 2. The effective Hamaker constant is proportional to the difference in the polarizabilities of the dispersed particles and the suspension medium:  $A_{off} = (\sqrt{A_{DM}} - \sqrt{A_{PAST}})^2$ , where  $A_{DM}$  and  $A_{PART}$  are the Hamaker constants for the suspension medium and the particles, respectively. As the suspended particles and the dispersion medium become similar in nature,  $A_{DM}$  and  $A_{PAST}$  become closer in magnitude, and  $A_{off}$  and  $V_A$  become unable. That is, by reducing the differences between the Hamaker constant associated with suspension medium and the Hamaker constant associated with the dispersed particles, the effective Hamaker constant (and corresponding van dar Weals attractive forces) may be reduced.

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One way to minimize the differences in the Hamsker constants is to create a "homodispersion", that is make both the continuous and dispersed phases assantially indistinguishable as discussed above. Benides exploiting the morphology of the particles to reduce the effective Hamsker constant, the components of the structural matrix (defining the perforated microstructures) will praferably be chosen so as to axhibit a Hamsker constant relatively close to that of the selected suspension medium. In this respect, one may use the actual values of the Hamsker constants of the suspension medium and the particulate components to determine the composibility of the dispersion ingredients and provide a good indication as to the stability of the preparation. Alternatively, one could select relatively compatible perforated microstructure components and suspension mediums using characteristic physical values that coincide with measurable Hamsker constants but are more readily discormible.

In this respect, it has been found that the refractive index values of many compounds tend to scale with the corresponding Hamaker constant. Accordingly, easily measurable refractive index values may be used to provide a fairly good indication as to which combination of suspension medium and particle excipients will provide a dispersion having a relatively low effective Hamaker constant and associated stability. It will be appreciated that, since refractive indices of compounds are videly evailable or easily derived, the use of such values allows for the formation of stabilized dispersions in accordance with the present invention without undue experimentation. For the purpose of illustration only, the refractive indices of several commends commentified with the disclosed dispersions are provided in Table Immediately below:

	Compound	Refractiva Index
	HFA-134a	1.172
	HFA-227	1.223
5	CFC-12	1.287
-	CFC-114	1.288
	PFOB	1.305
	Mannitol	1,333
	Ethanol	1.361
10 .	n-octane	1.397
	DMPC	1.43
	Pluronic F-68	1.43
	Sucrose	1.538
	Hydroxyathylstarch	1.54
15	Sodium chloride	1.544

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Consistent with the competible dispersion components set forth above, those skilled in the art will appreciate that, the formation of dispersions wherein the components have a refrective index differential of less than about 0.5 is preferred. That is, the refractive index of the suspension medium will preferably be within about 0.5 of the refractive index associated with the perforated particles or microstructures. It will further be appreciated that, the refractive index of the suspension medium and the particles may be measured directly or approximated using the refractive index of the major component in each respective phase. For the suspension medium, the major component way be determined on a weight percent besis. For the suspension medium, the major component will typically be derived on a volume percentage basis. In selected ambodiments of the present invention the refractive index differential value will prefarably be less than about 0.45, about 0.4, about 0.35 or evan less than about 0.3. Given that lower refractive index differentials of less than about 0.28, about 0.4, about 0.25 or evan less than about 0.3. Given that lower refractive index differentials of less than about 0.28, about 0.40, about 0.50 or evan less than about 0.50 or even less than about 0.50 o

As discussed above, the minimization of density differences between the particles and the continuous phase is largely dependent on the perforated and/or hollow nature of the microstructures, such that the suspension medium constitutes most of the particle volume. As used herein, the term "particle volume" corresponds to the volume of suspension medium that would be displaced by the incorporated hollow/porous perticles if they were solid, i.e. the volume defined by the particle boundary. For the purposes of explenation, and as discussed above, these fluid filled particulate volumes may be referred to as "virtual particles." Preferably, the avarage volume of the bioactive apentlescipient shall or matrix (i.e. tha volume of medium actually displaced by the perforated microstructure) comprises less than 70% of the virtual particle. More preferably, the volume of the microparticulate matrix

comprises less than about 60%, 40%, 30% or even 20% of the average particle volume. Even more praterably, the average avolume of the shallmentrix comprises less than about 10%, 5%, 3% or 1% of the average particle volume. Those skilled in the art will appreciate that, such a matrix or shell volumes typically contributes little to the virtual particle density which is overwhelmingly dictated by the suspension medium found therein. Of course, in selected embodiments the excipients used to form the perforated microstructure may be chosen so the density of the resulting matrix or shell approximates the density of the surrounding

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It will further be appreciated that, the use of such microstructures will allow the apparent density of the virtual particles to appreach that of the suspension medium substantially eliminating the attractive van der Weals forces. Moreover, as previously discussed, the camponants of the microparticulate matrix are preferably selected, as much as possible given other considerations, to approximate the density of suspension medium. Accordingly, in preferred embodiments of the present invention, the virtual particles and suspension medium will have a density differential of less than about 0.8 g/cm<sup>2</sup>. That is, the mean density of the virtual particles and less than about 0.8 g/cm<sup>2</sup> of the suspension medium. More preferably, the mean density of the virtual particles will be within 0.5, 0.4, 0.3 or 0.2 g/cm<sup>2</sup> of the selected suspension medium. In even more preferable embodiments the density differential will be less than about 0.1, 0.05, 0.01, or even less than 0.005 g/cm<sup>2</sup>.

In addition to the aforementioned edvantages, the use of hollow, porous particles allows for the formation of free-flowing dispersions comprising much higher volume fractions of particles in suspension. It should be appreciated that, the formulation of prior art dispersions at volume fractions approaching place packing generally results in drematic increases in dispersion viscoelestic behavior. Rheadogical behavior of this type is not appropriate for MOI applications. Those skilled in the art will appreciate that, the volume fraction of the particles may be defined as the ratio of the apparent volume of the particles (i.e. the particle volume) to the total volume of the system. Each system has a maximum packing fraction of 0.52 while those in a face centered cubic/hasagonal close packed configuration reach a maximum packing fraction of approximately 0.74. For non-spherical particles or polydisperse systems, the derived values are different. Accordingly, the maximum packing fraction is often considered to be an empirical parameter for a given system.

Here, it was supprisingly found that the porous structures of the present invancion do not exhibit undesirable viscoelastic bahavior even at high volume fractions, approaching does packing. To the contrary, they remain as free flowing, low viscosity suspensions having little or no yield stress whan compared with analogous suspensions comprising solid particulates. The low viscosity of the disclosed suspensions is thought to be due, at least in large part, to the relatively low van der Weals ettraction between tha fluid filled hollow, porous particles. As such, in selected ambodiments the volume fraction of the disclosed dispersions is

greater than approximately 0.3. Other embodiments may have packing values on the order of 0.3 to about 0.5 or on the order of 0.5 to about 0.8, with the higher values approaching a class packing condition. Moreover, as particle sedimentation tends to naturally decrease when the volume fraction approaches class packing, the formation of relatively concentrated dispersions may further increase formulation stability.

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Although the methods and compositions of the present invention may be used to form relatively concentrated suspensions, the stabilizing factors work equally well at much lower packing volumes and such dispersions are contemplated as being within the scope of the instant disclosure. In this regard, it will be appreciated that, dispersions comprising low volume fractions are extremely difficult to stabilize using prior at techniques. Conversely, dispersions incorporating perforated microstructures comprising a bisactive agent as described herein are particularly stable even at low volume fractions. Accordingly, the present invention allows for stabilized dispersions, and particularly respiratory dispersions, to the formed and used at volume fractions less than 0.3. In scome preferred embodiments, the volume fractions is approximately 0.0001 -0.3, more preferably 0.001 -0.01. Yet other preferred embodiments comprise stabilized suspensions having volume fractions from approximately 0.01 to approximately 0.1.

The perforated microstructures of the present invention may also be used to stabilize dilute suspensions of micronized bioscitive agents. In such embodiments the perforated microstructures may be added to increase the volume fraction of particles in the suspension, thereby increasing suspension stability to creaming or sedimentation. Further, in these embodiments the incorporated microstructures may also set in preventing closs approach (aggregation) of the micronized drug particles. It should be appreciated that, the perforated microstructures incorporated in such embodiments do not necessarily comprise a bioactive agent. Rather, they may be formed exclusively of various excipients, including surfactents.

Those skilled in the art will further appreciate that the stabilized suspensions or dispersions of the present invention may be prepared by dispersal of the microstructures in the selected suspension medium which may then be placed in a container or reservoir. In this regard, the stabilized preparedions of the present invention can be made by simply combining the components in sufficient quantity to produce the final desired dispersion concentration. Although the microstructures readily disperses without mechanical energy, the application of mechanical energy to aid in dispersion (e.g. with the aid of sonication) is contemplated, particularly for the formation of stable emulsions or reverse emulsions. Alternatively, the components may be mixed by simple shaking or other type of orgination. The process is preferably carried out under anhydrous conditions to advise any adverse effects of maisture on suspension stability. Once formed, the dispersion has a reduced susceptibility to floccation and sedimentation.

As indicated throughout the instant specification, the dispersions of the present invention are preferably stabilized. In a broad sense, the term "stabilized dispersion" will be held to mean any dispersion that resists aggregation, flocculation or creaming to the extent required to provide for the effective delivery of a biosoctive agent. While those skilled in the art will appreciate that there are severel methods that may be used to assess the stability

of a given dispersion, a preferred method for the purposes of the present invention comprises determination of creaming or sedimentation firm using a dynamic photosedimentation method. As seen in Example IX and Figure 2, a preferred method comprises subjecting suspended particles to a centrifugal force and emusing absorbance of the suspension as a function of time. A rapid decrease in the absorbance identifies a suspension with poor stability. It is submitted that these skilled in the art will be able to adapt the procedure to specific suspensions without undoe experimentation.

For the purposes of the present invention the craeming time shall be defined as the time for the suspended drug periodates to cream to 1/2 the volume of the suspended drug periodates to cream to 1/2 the volume as the suspension medium. Similarly, the sedimentation trim may be defined as the time it takes for the periodates the sediment in 1/2 the volume of the liquid medium. Besides the photosedimentation technique described above, a relatively simple way to determine the creaming time of a preparation is to provide the particulate suspension in a sealed glass vial. The visia are agitated or aboken to provide relatively homogeneous dispersions which are then set aside and observed using appropriate instrumentation or by visual inspection. The time necessary for the suspended periodates to cream to 1/2 the volume of the suspension medium (i.a., to rise to that top half of the suspension medium), or to sediment within 1/2 the volume (i.e., to settle in the bottom 1/2 of the medium), is then noted. Suspension formulations having a creaming time greater than 1 minute are preferred and indicates satisfalls stability. More preferredly, the stabilized dispensions comprise creaming times or preferred and indicates assistable stability. More preferred the preferred embodiments, the stabilized dispensions arithist creaming times of greater than ebout 1, 1.5, 2, 2.5, or 3 hours. Substantially equivalent puriods for sedimentation times are indicative of comentative discussions.

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As discussed herein, the stabilized dispersions disclosed harmin may preferably be administrated to the naseal or palmonary air passages of a patient the exercisation, such as with a material dose inhaler. The use of such stabilized preparations provides for superior dose reproducibility and improved lung deposition as described above. MOIs an well known in the art and could easily be employed for administration of the claimed dispersions without under experimentation. Breath activated MOIs, as well as these compassing other types of improvements which have been, or will be, developed are also compatible with the stabilized dispersions and present invention and, as such, are contemplated as being with in the scepa thereof. However, it should be emphasized that, in performed embodiments, this stabilized dispersions may be administrated with an MOI using a number of different mostes including, but not limited to, topical, assal, pulmoneny or one! Those skilled in the art will appreciate that, such routes are well known and that the dosing and administration procedures may be easily derived for the stabilized dispersions of the present invention.

MDI caristers generally comprise a container or reservoir capable of withstanding the vapor pressure of the propellant used such as, a plastic or plastic-costed glass buttle, or preferrity, a metal can or, for example, an aluminum can which may optionally be anodized, lacquer-ceeted and/or plastic-costed, wherein the contained closed with a metering valve. The metering valves are designed to definer a metered amount of the formulation per actuation. The valves incorporate a gasket to prevent leakage of propellant through the valve. The gasket may

comprise any suitable elastomeric meterial such as, for example, low density polyethylene, chlorobutyl, black and white butadiene-ecryloritrile nubbars, butyl rubber and neoprene. Suitable velvas are commercially available from manufacturars well known in the seresol industry, for example, from Valois, France (e.g. DFID, DF30, 0F31/50 ACT, DF60, Bespak plc, LTX (e.g. BK300, BK556) and 3M-Neotechnic Ltd, UK (e.g. Spraymiser).

Each filled center is conveniently fitted into a satistile channeling device or actuator prior to use to form a materned dose inhaler for administration of the medicament into the lungs or nasal cavity of a partient. Suitable channeling devices comprise for example a valve actuator and a cyfundrical or concilite passages through which medicament may be delivered from the filled canister via the metering valve, to the nose or month of a patient e.g., a mouthpiece actuator. Metered dose inhalers are designed to deliver a fixed unit dosage of medicament per actuation such as, for example, in the range of 10 to 5000 micrograms of bioactive egent per actuation. Typically, a single channel canister will covide for tens or even hundreds of shots or doses.

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With respect to MDIs, it is an advantage of the present invention that any biocompatible suspension medium having decaute vapor pressure to act as a propellant may be used. Particularly preferred suspension medium are compatible with use in a metered dose inhaler. That is, they will be able to form aerosals upon the activation of the metering valve and associated release of pressure. In general, the selected suspension medium should be biocompatible (i.e. reletively non-toxic) and non-reactive with respect to the suspension fur-formed microstructures comprising the bioactive agent. Preferably, the suspension medium will not act as a substantial solvent for any compenents incorporated in the perforated microstructures. Selected embodiments of the invention comprise suspension media selected from the group consisting of fluorocarbons (including those substituted with other hidogens), hydrocarbons, alcohols, at laters or combinations thereof. It will be appreciated thet, the suspension medium may comprise a mixture of various compounds selected to impart specific characteristics.

stratospheric ozone. In particular it is desirable that the formulations are substantially free of chlorofluorocarbons such as CCI<sub>2</sub>F, CCI<sub>2</sub>F<sub>2</sub>, and CF<sub>2</sub>CCI<sub>2</sub>.

Specific fluorecurbons, or classes of fluorinated compounds, that are useful in the suspension media include, but are not limited to, fluorohystame, fluorocycloheptame, fluoromethylcycloheptame, fluoromethylcycloheptame, fluoromethylcycloperature, fluorocycloperature, fluoromethylcycloperature, fluoromethylcycloperat

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In addition to the aforementioned fluorocarbons and hydrofluorocalkanes, various chlorofluorocarbons and substituted fluorinested compounds may also be used as suspansion mediums in accordance with the teachings herein. In this respect, FC-11 (CCL3R, FC-1181 (CBCL3R), FC-1182 (CBC2CFF, FC-1282 (CFC2R), FC-1282 (CHC2R), FC-1282 (CHC2R), FC-1282 (CHC3CF), FC-1282 (CHC3CF), FC-1282 (CHC3CF), FC-1282 (CHC3CF), FC-1282 (CHC3CF), FC-133 (CHC1FCHCI), FC-133 (CHC1FCHCI), FC-141 (CHC2CHCHCI), FC-1418 (CC1FCHCI), FC-142 (CHC2CHCCI), FC-157 (CHC2CHCHCI), FC-1121 (CHC1-CFCI), FC-11

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Along with the aforementioned embodiments, the stabilized dispersions of the present invention may also be used in conjunction with nebulizers to provide an aerosolized medicament that may be administared to the pulmonery eir passages of a patient in need thereof. Nebulizers are well known in the art and could easily employed for administration of the claimed dispersions without undue experimentation. Breath activated nebulizers, as well as those comprising other types of improvements which have been, or will be, developed are also computable with the stabilized dispersions and present invention and are contemplated as being within the scope thereof.

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Nebulizers work by forming aerosols, that is converting a bulk liquid into small droplets suspended in a breathable gas. Here, the aerosolized medicament to be administered (preferably to the pulmonary air passages) will comprise small droplets of suspension medium associated with perforated microstructures comprising a bioactive agent. In such embodiments, the stabilized despersions of the present invention will typically be placed in a fluid reservoir operably associated with a nebulizer. The specific volumes of preparation provided, means of filing the reservoir, etc., will largely be dependent on the selection of the infinitional nebulizer and is well writhin the purview of the skilled artisen. Of course, the present invention is entirely compatible with single-dose nebulizers and multiple free nebulizers.

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Traditional prior art nabulizer preparations typically comprise aqueous solutions of the selected pharmaceutical compound. With such prior art nebulizer preparations, it has long been established that corruption of

the incorporated therapeutic compound can severely reduce efficacy. For example, with conventional equeous multidose nebulizer preparations, bectain contentination is a constant problem. In addition, the solubilized medicarent may precipitate out, or degrade over time, adversely effecting the delivery profile. This is particularly true of larger, may lead to particle growth that results in a substantial and reduction in lung penetration and a corresponding decrease in bioavailability. Such decira inconnomics marked vecesses the effectiveness of any treatment.

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The present invention overcomes these and other difficulties by providing stabilized dispersions with a suspension medium that prelarably comprises a fluorinated compound (i.e. a fluorochemical, fluorocarbon or perfluorocarbon). Perticularly preferred embodiments of the present invention complises fluorochemicals that are liquid at room temperstum. As indicated above, the use of such compounds, whether as a continuous phase or, es a suspension medium, provides several advantages over prior art liquid inhaldion preparations. In this regard, it is well established that many fluorochemicals have a proven history of safety and blocompetibility in the lung. Further, in contrast to aqueous solutions, fluorochemicals do not negatively impact gas exchange following pulmonary administration. To the contrary, they may actually be able to improve gas exchange and, due to their unique wettability characteristics, are able to carry an aerosotized stream of particles deeper into the lung, thereby improving systemic delivery of the desired phaseoutical compound. In addition, the relatively non-reactive neture of fluorochemicals acts to retard any degradation of an incorporated bisocitive agent. Finally, many fluorochemicals and substitute their processing the potential for microbial crowthin competible nebulizer devices.

In any event, rebufazer mediated aerosolization typically requires an input of energy in order to produce the increased surface area of the droplets and, in some cases, to provide transportation of the atomized or aerosolized medicament. One common mode of aerosolization is forcing a stream of fluid to be ejected from a nozzle, whereby droplets are formed. With respect to nebufaze deministration, additional energy is usually imparted to provide droplets that will be sufficiently small to be transported deep into the lungs. Thus, additional energy is needed, such as that provided by a high velocity gas stream or a piezoelectric crystal. Two popular types of nebufazes, jet nebufazes and ultrazoric nebufazes, rely on the aforementioned methods of applying additional energy to the fluid during atomization.

In terms of pulmonery delivery of bioactive agents to the systemic circulation via nebulization, recent research has focused on the use of portable hand-held ultrasseic nebulizars, also referred to as metered solutions. These devices, generally known as single-bolus nebulizers, sensatize a single-bolus of medication in an equeous solution with a particle size efficient for feep lung delivery in one or two breaths. These devices fall into three broad categories. The first category comprises pure piscodectric single-bolus nebulizers such as those described by Mütterlein, et. al., U. Aerosal Med. 1988; 12311. In another category, the desired aerosal cloud may be generated by microchannel extrusion single-bolus nebulizers such as those described in U.S. Pat. No. 3,812,654. Finally, a third category comprises devices exemptified by Robertson, et. al., (WO 82/11050) which describes cyclic pressurization single-bolus nebulizers. Each of the afortementomed references is incorporated herein in their antitry. Med.t devices

are manually actuated, but some devices exist which are breath actuated. Breath actuated devices work by releasing acrossor when the device sanses the patient inheling through a circuit. Breath actuated nebulizers may also be placed in-like on a ventilator circuit to release serosoil into the air flow which comprises the inspiration gases for a nation.

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Regardless of which type of nebufaer is employed, it is an advantage of the present invention that biocompatible nonequeous extremounts may be used as suspension mediums. Preferably, they will be able to form services upon the application of anergy therets. In general, the selected suspension medium should be biocompatible (i.e. relatively non-toxic) and non-reactive with respect to the suspenside perforated microstructures compising the bioactive agent. Preferred embodiments comprise suspension medias selected from the group consisting of flourochemicals, fluanceathous finduling those substituted with other halogous's perfluorocarbons, fluorocerbon/hydrocation dilocks, hydrocarbons, alcohold, ethers, or combinations thereof. It will be appreciated that, the suspension medium may comprise a mixture of various compounds selected to impart specific cheracteristics. It will also be appreciated that the perforated microstructures are preferrably insoluble in the suspension medium, thereby providing for stabilized medicament particles, and effectively protecting a selected bioactive agent from degradation, as might occur during prolonged storage in an aqueous solution. In preferred embodiments, the selected suspension medium is bacteriotetic. The suspension formulation also protects the bioactive agent from degradation during the nebufazation process.

As indicated above, the suspension media may comprise any one of a number of different compounds including hydrocarbons, fluorocarbons or hydrocarbons or hydrocarbons, fluorocarbons or hydrocarbons or highly fluorinated or perfluorocarbon disposal may be lines; branched or cyclic, saturated or unsaturated compounds. Conventional structural derivatives of these fluorochamicals and hydrocarbons are also contemplated as being within the scope of the present invention as well. Selected embodiments comprising these totally or partially fluorinated compounds may contain one or more hetero-atoms and/or atoms of branine or chlorine. Preferably, these fluorochamicals comprise from 2 to 16 carbon atoms and include, but are not limited to, linear, cyclic or polycyclic perfluoroal/kanes, biologerfluoroal/kylakinese, perfluorochemse, perfluoroal/kanes, bringerfluoroal/kylakinese, perfluorochemse, perfluoroal/kylakinese, perfluoroal/kylakines

More generally, exemplary fluorechamicals which are contemplated for use in the present invention generally include halogeneted fluorechamicals (i.e.  $C_{F_m} \times X$ ,  $KC_{F_m} X$ , where  $n = 2\cdot 10$ , X = Br,  $Cl \ or \ ll and, in particular, 1-bramo-Factane <math>n \cdot C_{F_m} Br$ , 1-bramo-Factane  $n \cdot C_{F_m} Br$ , 1-bramo-Factane in  $C_{F_m} Br$ , 1-bramo-Factane in  $C_{F_m} Br$ , 1-bramo-Factane in U.S. and 1.8-distance from the second in U.S. Patent No. 3,975.512 to long and are incorporated harbin by reference. Specific fluorechamicals having chloride substituents.

such as perfluoreoctyl chloride (n-C<sub>0</sub>F<sub>12</sub>Cl), 1,8-dichlore-F-octane (n-ClC<sub>2</sub>F<sub>16</sub>Cl), 1,6-dichlore-F-hexane (n-ClC<sub>2</sub>F<sub>12</sub>Cl), and 1. 4-dichlore-F-hutane (n-ClC,F,Cl) are also preferred.

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Polycyclic and cyclic fluorochemicals, such as C<sub>P</sub>F<sub>11</sub> (F-decalin or parluorodecalin), perfluorophenanthrena, perfluorodetermenthytrycholexame (AP-144) and perfluoro abuytdecalin are also within the scope of the invention. Additional useful fluorochemicals include perfluorosted armines, such as F-triproplamine ("FTPA") and F-tributylamine ("FTBA"). F-4-methylocalinydroquinoidine ("FTMO"). F-1N-methylocalinydroquinoidine ("FTMO"). F-1N-methyloca

Specific fluorocarbons, or classes of fluorinated compounds, that may be useful as suspension media include, but are not limited to, fluorobystame, fluorocyclohegrane fluoromethylcyclohegrane, fluorom

While any fluid compound capable of producing an sereard upon the application of energy may be used in conjunction with the present invention, the selected suspension medium will preferrely have a vapor pressure less than about 5 atmospheres and more preferably less than about 2 atmospheres. Unless otherwise specified, all vapor pressures not extend therein are measured at 25°C. In other embediaments, preferred suspension media compounds will have vapor pressures on the order of about 5 terr to about 700 terr, with more preferable compounds will have vapor pressures on the order of from about 8 terr to about 500 terr, while still more preferable compounds will have vapor pressures on the order of from about 10 terr to about 350 torr. Such suspension media may be used in conjunction with compressed sir nebulizers, ultrasoric nebulizers or with mechanical atomizers to provide effective ventiletion therepy. Moreover, more volatic compounds may be mixed with lower vapor pressure components to provide suspension media having specified physical characteristics selected to further improve stability or enhance the biosvellability of the disposated biasticine asant.

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Other embodiments of the present invention directed to netulizers will comprise suspension media that bail at selected temperatures under ambient conditions. Se. 1 atml. For example, preferred embodiments will comprise suspension media compounds that boil above 0°C, above 5°C, above 10°C, done 15°, or above 20°C. In other ambodiments, the suspension media compound may boil at or above 25°C or at or above 30°C. In yet other ambodiments, the selected suspension media compound may boil at or above human body temperature (i.e. 37°C), above 45°C, 55°C, 65°C, 75°C, 65°C or above 100°C.

Along with MOIs and nebulizers, it will be appreciated that the stabilized dispersions of the present invention may be used in conjunction with fiquid dose instillation or LOI techniques. Liquid dose instillation involves the direct administration of a stabilized dispersion to the lung. In this regard, direct pulmonary administration of bloactive compounds is particularly affective in the treatment of disorders aspecially where poor vascular circulation of diseased portions of a lung reduces the effectiveness of intrevenous drug delivery. With respect to LOI the stabilized dispersions are preferably used in conjunction with partial liquid ventilation or total liquid ventilation. Moreover, the present invention may further comprise introducing a therapeutically beneficial amount of a physiologically acceptable gas (such as ritric acids or oxygen) into the pharmaceutical microdispersion prior to, during or following administration.

For LDI, the dispersions of the present invention may be administered to the lang using a pulmonary delivery conduit. Those skilled in the art will appreciate the term "pulmonary delivery conduit", as used herein, shall be construed in a broad sense to comprise any device or apparatus, or component thereof, that provides for the instillation or administration of a liquid in the lungs. In this respect a pulmonary delivery conduit or delivery conduit shall be held to mean any bone, luman, catheter, tube, conduit, syringe, actuator, mouthpiece, and addreched tube or bronchoscope that provides for the administration or instillation of the disclosed dispersions to at least a portion of the pulmonary air passages of a patient in need thereof. It will be appreciated that the delivery conduit may or may not be associated with a liquid ventilator or gas ventilator.

In particularly preferred embodiments the delivery conduit shall comprise an endotracheal tube or bronchoscope.

Here it must be emphasized that the dispersions of the present invention may be administered to ventilated (e.g. those connected to a mechanical ventilater) or nonventilated, patients (e.g. those undergoing spontaneous respiration). Accordingly, in preferred embodiments the methods and systems of the present invention may comprise the use or inclusion of a mechanical ventilater. Further, the stabilized dispersions of the present invention may also be used as a lavage agent to remove debris in the lung, or for diagnostic lavage procedures. In eny case the introduction of liquids, perticularly fluorochemicals, into the lungs of a patient is well known and could be accomplished by a skilled artisen in passession of the instant specification without under experimentation.

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Those skilled in the ent will appreciate that suspension media compatible with 101 techniques are similar to thase set forth above for use in conjunction with nebotizers. Accordingly, for the purposes of the present application suspension media for dispersions compatible with LOI shall be equivalent to those enumerated above in conjunction with use in nebulizers. In any event, it will be appreciated that in particularly preferred LOI embodiments the selected suspension medium shall comprise a fluorochemical that is finial under emblent conditions.

It will be understood thet, in connection with the present invention, the disclosed dispersions are preferably administered directly to at least a portion of the pulmonary air passages of a mammal. As used herein, the terms "direct instillation" or "direct administration" shall be held to mean the introduction of a stabilized dispersion into the lung cevity of a mammal. That is, the dispersion will preferably be edministered through the traches of a patient end into the lungs as a relatively free flowing fluid passing through a delivery conduit end into the pulmonery air passages. In this regard, the flow of the dispersion may be grevity assisted or may be afforded by induced pressure such as through a pump or the compression of a syringe plunger. In any case, the amount of dispersion administered may be monitored by mechanical devices such as flow meters or by visual inspection.

While the stabilized dispersions may be administered up to the functional residual capacity of the lungs of a patient, it will be appreciated that selected embodiments will comprise the pulmonary administration of much smaller volumes (e.g. on the order of a milliter or less). For example, depending on the disorder to be treated, the volume administrated may be on the order of 1, 3, 5, 10, 20, 50, 100, 200 or 500 milliters. In preferred embodiments the liquid volume is less than 0.25 or 0.5 percent FRC. For particularly preferred embodiments, the liquid volume is 0.1 percent FRC or less. With respect to the administration of relatively low volumes of stabilized dispersions it will be appreciated that the veretakility and spreading characteristics of the suspension media (particularly fluorechemicals) will facilitate the even distribution of the bioactive agent in the lung. However, in other embodiments it may be preferable to administer the suspensions a volumes of greater than 0.5, 0.75 or 0.9 parcant FRC. In any event. Did treatment as disclosed herein expressors a new atternative for critically

ill patients on mechanical ventiletors, and opens the door for treatment of less ill patients with bronchoscopic administration.

It will also be understood that other components can be included in the stabilized dispersions of the present invention. For example, comotic agents, stabilizers, cheletors, buffers, viscosity modiators, saits, and sugaes can be added to fine turne the stabilized dispersions for maximum file and asse of administration. Such components may be added directly to the suspension medium or associated with, or incorporated in, the perforated microstructures. Considerations such as stability, isotonicity, and biocomposibility may govern the use of commonland additives to the discisced compositions. The use of such agents will be understood to those of ordinary skill in the art and, the specific quantities, ratios, and types of agents can be determined empirically without under experimentation.

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Moreover, while the stabilized dispersions of the present invention are particularly suitable for the pulmonary administration of bioactive agents, they may also be used for the localized or systemic administration of compounds to any location of the body. Accordingly, it shedd be emphasized that, in preferred embodiments, the formutations may be administrated using a number of different routes including, but not limited to, the gestrointessinal tract, the respiratory tract, topically, intramucadarly, intrapentaneally, assally, vaginally, rectally, surelly, routly or outset. More generally, the stabilized dispersions of the present invention may be used to deliver agents topically or by administration to e non-pulmonary body cavity. In preferred embodiments the body cavity is selected from the group consisting of the peritoneum, sinus cavity, rectum, urethra, gastrointestinal tract, nased cavity, vagina, auditory mentus, and cavity, boccal pouch and pleura. Among other indications, stabilized dispersions comprising the appropriate bioactive eyem, (e.g., an antibiotic or an anti-inflammatory), may be used to treat infections of the eye, sinustis, infections of the auditory tract and even infections or disorders of the gestrointestinal tract. With respect to the latter, the dispersions of the present invention may be used to selectively deliver pharmaceutical compounds to the latter, the dispersions of the tractment of H<sub>c</sub> power/infections or other uteer related disorders.

With regard to the perforated microstructure powders and stabilized dispersions disclosed harein those skilled in the art will appreciate that they may be advantageously supplied to the physician or other health perforasional, in a starile, prepeckaged or kit form. More particularly, the formulations may be supplied as stable powders or preformed dispersions ready for administration to the patient. Conversely, they may be provided as separate, ready to mix components. When provided in a ready to use form, the powders or dispersions may be packaged in single use containers or reservoirs, as well as in multi-use containers or reservoirs. In either case, the container or reservoir may be associated with the selected inhelation or administration device and used as described herein. When provided as individual components (e.g., as providered microspheres and as neal suspension medium) the stabilized preparations may then be formed at any time prior to use by simply combining the contents of the containers as directed. Additionally, such kits may contain a number of ready to mix, or prepackaged dosing units so that the user can then administer tham as needed.

Although preferred embodiments of the present invention comprise providers and stabilized dispersions for use in pharmaceutical applications, it will be approached that the perforated microestructures and disclosed dispersions may be used for a number of non pharmaceutical applications. That is, the present invention provides perforated microestructures which have a broad range of applications where a powder is suspended endlor sensatized. In particular, the present invention is expecially affective where an active or bioective ingredient must be dissolved, suspended or solubilized as fast as possible. By increasing the surface area of the porous microparticles or by incorporation with suitable accipients as described herein, will result in an improvement in dispersibility, and/or suspension stability. In this repard, rapid dispersement applications include, but are not limited to: detergents, dishwasher detergents, food sweeteners, condiments, spices, mineral flotation detergents, thickering agents, folier fertikers, phytohermones, insect pheromones, insect repellents, pert receillents, pesticides, functiones, disinfecturnts, perfunes, deadcrants, atc.

Applications that require finely divided particles in a non-equeous suspension media (i.e., solid, liquid or gessous) are also contemplated as being within the scope of the present invention. As explained herein, the use of perforated microstructures to provide a "homodispersion" minimizes particle-particle interactions. As such, the perforated microspheres and stabilized suspensions of the present invention are particularly compatible with applications that require: inorpacin pigments, dyes, inks, paints, explasives, pyratechnic, addordnants, absorbents, absorbents, absorbents, absorbents, absorbents, between the propertions of the present invention offers benefits over prior art preparations for use in applications which require serosolization attainization. In such non-pharmaceusical uses the preparations can be in the form of a liquid suspension (such as with a propellant) or as a dry powder. Preferred embodiments comprising perforated microstructures as described herein include, but are not limited to, ink jet printing formulations, powder coating, spray paint, spray-pesticides etc.

The foregoing description will be more fully understood with reference to the following Examples. Such Examples, are, however, merely representative of preferred methods of practicing the present evention and should not be read as limiting the scope of the invention.

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#### Preparation of Hollow Porous Particles of Gentamicin Sulfata by Spray Drying

40 to 60ml of the following solutions were prepared for spray drying:

30 50% w/w hydrogeneted phosphetidylcholine, E-100-3 (Lippid K.G., Ludwigshefen, Germeny) 50% w/w gentemicin sulfate (Amresco, Selon, OH) Perfluorocctylbromide, Perflubron (NMK, Japan) Deionized water

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Perforated microstructures comprising gentamicin sulfate were prepared by a spray drying technique using a B-191 Mini Spray-Drier (Büchi, Hawil, Switzerland) under the following conditions:

aspiration: 100%, inlet temperature: 85°C; outlet temperature: 61°C; feed pump: 10%;  $N_2$  flow: 2,800 L/hr. Vaniations in powder porosity were examined as a function of the blowing agent concentration.

Fluorocarboe-in-water emulsions of perfluorocatyl bromide conteiring a 1:1 w/w ratio of phesphathylcholine (PC), and gentamicin additate ware prepared varing only the PFC/PC ratio. 1.3 grams of hydrogenated egg phosphathylcholine was dispersed in 25 nt. deionized water using a Ultra-Turrax mixer model 1:251 at 8000 rym for 2 to 5 minutes (T = 60-70°C). A range from 0 to 40 grams of perfluithorn was added drowpxise during mixing (T = 60-70°C). After addition was complete, the fluorocarbon-in-water amulsion was mixed for an additional period of not less than 4 minutes. The resulting coerse emulsions were then homogenized under high pressure with an Avestin (Ottawa, Canada) homogenizer at 15,000 psi for 5 passes. Gentamicin suffate was dissolved in approximately 4 to 5 mt. deionized water and subsequently mixed with the perfluition emulsion immediately prior to the spray dry process. The gentamicin powders were then obtained by spray drying using the conditions described above. A free flowing pals yellow prowder was obtained for all perfluitron containing formulations. The yield for each of the various formulations ranged from 35% to 60%.

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## Morphology of Gentamicin Sulfate Spray-Dried Powders

A strong dependence of the powder morphology, degree of perceity, and production yield was observed as a function of the PFC/PC ratio by scanning electron microscopy (SEM). A series of six SEM micrographs illustrating these observations, labeled 1A1 to 1FI, are shown in the left hand column of Fig. 1. As seen in these micrographs, the porcestly and surface roughness was found to be highly dependent on the concentration of the blowing agent, where the surface roughness, number and size of the pores increased with increasing PFC/PC ratios. Fer example, the formulation devoid of perfluorocctyl brande produced microstructures that appeared to be highly agglomerated and readily adhered to the surface of the glass vial. Similarly, smooth, spherically shaped microparticles were obtained when relatively little (PFC/PC ratio - 1.1 or 2.2) blowing agent was used. As the PFC/PC ratio was increased dramatically.

As shown in the right hand column of Fig. 1, the hollow nature of the microstructures was also enhanced by the incorporation of additional blowing agent. More particularly, the series of six micrographs labeled 1A2 to 1F2 show cross sections of fractured microstructures as revealed by transmission electron microscopy (TEM). Each of these images was produced using the same microstructure preparation as was used to produce the corresponding SEM micrograph in the left hand column. Both the hellow nature and wall thickness of the resulting perforated microstructures appeared to be largely dependent on the concentration of the selected blowing agent. That is, the hollow nature of the preparation appeared to increase and the hickness of the perticle walls appeared to electess as the PEC/PC ratio increased. As may be seen in Figs. 1A2 to 102 substantially said structures were obtained from formulations containing little or no fluorocathon blowing

agent. Conversely, the perforated microstructures produced using a relatively high PFC IPC ratio of approximately 45 (shown in Fig. 1F2 proved to be extremely hollow with a relatively thin wall ranging from about 43.5 to 26 from. Both types of particles are competible for use in the present invention.

## <u>Preparation of Spray Oried Gentamicin</u> Sulfate Particles using Various Blowing Agents

40 milliliters of the following solutions were prepared for spray drying:

40 milliliters of the following solutions were prepared for spray drying

50% w/w Hydrogenated Phosphatidylcholine, E100-3 10 (Lipoid KG, Ludwigshafen, Germanyl

> 50% w/w Gentamicin Sulfate (Amresco, Solon Ohio) Deionized water.

Blowing Agents:

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Parfluorodecalin, FDC (Air products, Allanton PA)
Parfluorooctylbromide, Parflubron (Atochem, Paris, France)
Parfluorhaxana, PFH (3M, St. Paul, MI)
1,12-trichlorottifluoroethane, Freon 113 (Baxter, McGaw Park, IL)

Hollow porous microspheres with a model hydrophilic drug, e.g., gernamicin sulfate, were prepared by spray drying. The blowing agent in thase formulations consisted of an amulating fluorochemical (FC) oil. Emulsions were prepared with the following FCs: PHI, Freon 113, Perflubtron BCc. 1.3 grams of the hydrogenated egg phosphatidylcholine was dispensed in 25 mL deionized waster using a Ultre-Turrax mixer immodel 7:28j at 81000 rpm for 2 to 5 minutes 11 - 60-70, 25 grams of FC was added dropwise during mixing (T - 60-70°C). After the addition was complete, the FC-in-waster mulsion was mixed for a total of not less than 4 minutes. The resulting emulsions were then further processed using an Avestin (Ottawa, Canada) high pressure homoganizer at 15,000 psi and 5 passes. Gentamicin sulfate was dissolved in approximately 4 to 5 mL deionized water and subsequently mixed with the FC emulsion. The gentamicin providers were obtained by spray drying (Blichi, 191 Mini Spray Dryer). Each emulsion was fed at a rate of 2.5 mL/min. The inlat and outlist temperatures of the spray dryer were 85°C and 55°C respectively. The nebalization air and aspiration flows were 2800 U.Hn and 100% respectively.

A free flowing pale yellow dry powder was obtained for all formulations. The yield for the various formulations ranged from 35 to 80%. The various gentamicin sulfate powders had a mean volume weighted particle diameters that ranged from 1.52 to 4.91 µm.

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IV

<u>Effect of Blowing Agent on the Morphology of</u>

Gentamicin Sulfata Spray-Driad Powders

A strong dependance of the powder morphology, poresity, and production yield (amount of powder captured in the cyclone) was observed as a function of the blowing agent building point. In this respect the powders produced in Example III were observed using assuring electron microscopy. Spray drying a fluorochemical (F1) emulsion with a boiling point below the 55°C outlet temperature (e.g., perfluorobexane IPFH) or Freen 1131, yielded amorphously shaped (shriveled or deflated) powders that contained little or no pores. Wherasa, emulsions formulated with higher boiling FCs (e.g., perfluorob, perfluorodeccialin, FDC) produced spherical powders practicles. Powders produced with higher boiling blowing agents also had production yields approximately two times greater than powders produced using relatively low boiling point blowing agents. The selected blowing agents also had production grants.

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Table II

Blowing Agent	(bp °C)
Freon 113	47.6
PFH	56
FDC	141
Perflubron	141

Example IV illustrates that the physical characteristics of the blowing agent (i.e., boiling point) greatly influences the ability to provide perforeted microparticles. A particular advantage of the present invention is the ability to after the microstructure morphology and porosity by modifying the conditions and nature of the blowing spent.

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## Preparation of Spray Dried Albuterol Sulfate Particles using Various Blowing Agents

Approximately 185 ml of the following solutions were prepared for spray drying:

49% w/w Hydrogenated Phosphatidylcholine, E100-3
(Lippid KG, Ludwigshafen, Germany)

50% w/w Albuteral Sulfate

(Accurate Chemical, Westbury, NY)

1% w/w Poloxamer 188, NF grade (Mount Olive, NJ) Deignized water

30 Blowing Agents:

Perfluorodecalin, FDC (Air products, Allenton PA)

Perfluorooctylbromide, Perflubron (Atochem, Paris) Perfluorobutylethane F4H2 (F-Tech, Japan)

Perfluorotributylamine FTBA (3M, St. Paul, MN)

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Albuterol sulfate powdar was prepared by spray-drying technique by using a B-191 Mini Spray-Drier (Büchi, Flawil, Switzarland) under the following conditions:

Aspiration: 100%

Inlet temparatura: 85°C Outlat temperature: 61°C Faad pump: 2.5 mL/min. N<sub>2</sub> flow: 47 L/min.

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The feed solution was prepared by mixing solutions A and B prior to spray drying.

<u>Solution A</u>: Twenty grams of water was used to dissolva 1.0 grams of Albuterol sulfate and 0.021 grams of poloxemer 188.

Solution B represented an emulsion of a fluorocarbon in water, stabilized by a phospholipid, which was prepared in the following way. Hydrogenated phosphatidylchinine (1.0 grama) twas homogenized in 130 grams of hot deionized water (T = 50 to 60°C) using an Ultra-Turax mixer (model T-25) at 8000 pr.m, for 2 to 5 minutes (T = 60.70°C). Twenty-five grams of Perfluoron (Atochen, Paria, France) was added dropwise during mixing. After the addition was complete, the Florochemical-in-water sentision was mixed for at least 4 minutes. The restifing emulsion was then processed using an Avestin (Ottava, Canada) high-pressure homogenizer at 18,000 psi and 5 passes. Solutions A and B were combined and fed into the spray dryer under the conditions described above. A free flowing, white powder was collected at the cyclene separator as is standard for this spray dryer. The albuterol saltest powders had mean volume valued particle diemeters ranging from 1.28 to 2.77 µm, as determined by an Aerosizer (Amherst Process Instruments, Amharst, MA). By SEM, the albutered saffetsiphospholipid spray dried powders were spherical and highly process.

Example V further demonstrates the wide variety of blowing agents that may be used to provide perforated microparticles. A particular advantage of the present invention is the ability to after the microstructure morphology and porosity by manipulating the formulation and spray drying conditions. Furthermore, Example V demonstrates the particle diversity achieved by the present invention and the ability to affectively incorporate a wide variety of pharmacousical agents therein.

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# VI Preparation of Hollow Porous PVA Particles by Spray Drying a Water-in-oil Emulsion

100 ml of the following solutions were prepared for spray drying:

80% w/w Bis-12-ethylheryll Sulfrasuccinic Sodium Salt,
(Aerosol OT, Kodak, Rochester, NY)
20% w/w Polyvinyl Alcohd, average malecular weight -30,000-70,000
(Sigma Chemicals, St. Louis, MO)
Carbon Tatrachloride (Aldrich Chemicals, Milwaukee, Wi)

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Aerosol OT/polyvinyl alcohol particles were prepared by spray-drying technique using a B-191 Mini Spray-Drier (Büchi, Flawii), Switzerland) under the following conditions:

Aspiration: 85%

Deionized water.

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#### WO 99/16419

Inlet temperature: 60°C Outlet temperature: 43°C Feed pump: 7.5 mL/min. N<sub>2</sub> flow: 36 L/min.

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Solution A: Twenty grams of water was used to dissolve 500 milligrams of polyvinyl alcohol (PVA).

Solution B: represented an emulsion of carbon tetrachloride in water, stabilized by aerosol OT, which was prepared in the following way. Two grams of serosol OT, was dispersed in 80 grams of carbon tetrachloride using a Ultre-Turrax mixer femded 1-25 et 8000 rpm for 2 to 5 minutes (T - 15° to 20°C). Twenty grams of 2.5% w/v PVA was added dropwise during mixing. After the addition was competet, the water-lineal emulsion was mixed for a total of not less than 4 minutes (T - 15° to 20°C). The resulting emulsion was then processed using an Avestin (Otteva, Canada) high-pressure hemogenizer at 12,000 psi and 2 passes. The emulsion was then fed into the spray driver under the conditions described above. A free flowing, white powder was celected at the cyclone seperator as is standard for this spray dryer. The Aerosol OTIPVA powder had a mean volume veighted particle diameter of 5.28 ± 3.27 µm as determined by an Aerosizer (Amherst Process Instruments, Amherst, MA).

Example VI further demonstrates the variety of emulsion systems (here, reverse wetar-in-oil), formulations and conditions that may be used to provide perforated microparticles. A particular advantage of the present invention is the ability to alter formulations and/or conditions to produce compositions having a microstructure with selected possity. This principle is further illustrated in the following example.

#### VΙΙ

# <u>Preparetion of Hollow Porous Polyceprolactone</u> Particles by Spray Drying a Water-in-Oil Emulsion

Particles by Spray Drying a Water-in-Oil Emulsis

25 100 mls of the following solutions were prepared for spray drying:

80% w/w Sorbitan Monostearate, Span 60 (Aldrich Chemicals, Milwaukee, WI)
20% w/w Polycaprolactone, average molecular weight = 65,000
(Aldrich Chemicals, Milwaukee, WI)

30 Carbon Tetrachloride (Aldrich Chemicals, Milwaukee, WI)
Descrized water.

Span 60/polycaprolactone particles were prepared by spray-drying technique by using a B-191 Mini Spray-Orier (Büchi, Rewil, Switzerland) under the following conditions:

35 Aspiration: 85% Inlet temperature: 50°C Outlet temperature: 38°C Fead pump: 7.5 mL/min. N. flow: 36 L/min.

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A watar-in-carbon tetrachloride emulsion was prepared in the following manner. Two grams of Spen 60, was dispersed in 80 grams of carbon tetrachloride using an Ultra-Turrax mizer Imodel T-25) at 8000 rpm for 2 to 5 minutes 1T – 15 to 20°C). Twenty grams of deionized watar was added dropwise during maining. After the addition was complete, the water-inoil emulsion was minutes for a total of not less than 4 minutes IT – 15 to 20°C). The resulting emulsion was then further processed using an Avestin (10 town, Canadal high-pressure hamogenizer at 12,000 psi and 2 passes. Five hundred milligrams of polycaprolectione was added directly to the amulsion and, mixed until thoroughly dissolved. The amulsion was then fed into the spray dryer under the conditions described above. A free Rewing, white powder was collected at the cyclone separator as is stendard for this dryer. The resulting Span 60(pplycaprolectione powder had a mean volume weighted particle diameter of 3.15 ± 2.17 µm. Again, the present Example demonstrates the versettlifty the instent invention with header to the feed stock used to provide the desired perforated microstructure.

VII

#### 15 Preparation of hollow porous powder by soray drying a casin-water emulsion

The following solutions were prepared with water for injection:

Solution 1:

3.9% w/v m-HES hydroxyethylsterch lAjinomoto, Tokyo, Jepenl 20 3.25% w/v Sodium chloride (Mallinckradt, St. Louis, MOI 2.83% w/v Sodium phospheta, dibasic (Mallinckradt, St. Louis, MOI 0.42% w/v Sodium phospheta, monobasic (Mallinckradt, St. Louis, MOI

Solution 2:

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0.45% w/v Poloxamer 188 (BASF, Mount Olive, NJ)
1.35% w/v Hydroganated egg phosphatidylcholine, EPC-3

|Lipoid KG, Ludwigshafen, Germany|

The ingredients of solution 1 were dissolved in warm water using a stir plate. The surfactants in solution 2 30 were dispussed in water using a high shear mixer. The solutions were combined following emulsification and saturated with infringen pint to spary dyring:

The resulting dry, free flowing, hollow spherical product had a mean particle diameter of  $2.6 \pm 1.5$   $\mu m$ . The particles were spherical and porous as determined by SEM.

This axemple illustrates the point that a wide of blowing agents there nitrogen may be used to provide microstructures axhibiting the desired morphology. Indeed, one of the primary advantages of the present invention is the ability to alter formation conditions so as to preserve biological activity (i.e. with protains), or to produce microstructures having selected poresity.

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## Suspension Stability of Gentamicin Sulfete Spray-Dried Powders

The suspension stability was defined as, the resistance of powders to cream in a nonequeous madium using a dynamic photosedimentation method. Each sample was suspanded in Perflubron at a concentration of 0.8 mg/tml.. The creaming rates were measured using a Horiba CAPA-700 photosedimentation particle size analyzer (Irvine, CA) under the following conditions:

> D (max): 3.00 m D (min.): 0.30 µm D (Div): 0.10 µm

3000 rpm

Rotor Speed: X: 10 mm 10

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The suspended particles were subjected to a centrifugal force and the absorbance of the suspension was measured as a function of time. A rapid decrease in the absorbance identifies a suspension with poor stability. Absorbance data was plotted versus time and the area under the curve was integrated between 0.1 and 1 min., which was taken as a relative measurament of stability. Figure 2 graphically depicts suspension stability as a function of PFC/PC ratio or porosity. In this case, the powder porosity was found to increase with increasing PFC/PC. Maximum suspension stability was observed with formulations having PFC/PC ratios between 3 to 15. For the most part, these formulations appeared stable for periods greater than 30 minutes using visual inspection techniques. At points beyond this ratio, the suspansions flocculated rapidly indicating decreased stability. Similar results were observed using the cream layer ratio method, where it was observed that suspansions with PFC/PC ratios batwaan 3 to 15 had a raduced craam layer thickness, indicating favorable suspension stability.

### Preparation of Hollow Porous Particles of Albuterol Sulfate by Spray-Drying

Hollow porous albutarol sulfata particles were prepared by a spray-drying tachnique with a B-191 Mini Spray-Drier (Büchi, Flawil, Switzerland) under the following spray conditions: aspiration: 100%, inlet temperature: 85°C; outlet temperature: 61°C; feed pump: 10%; N, flow: 2,800 L/hr. The feed solution was prepared by mixing two solutions A and B immediately prior to spray drying.

Solution A: 20g of water was used to dissolve 1g of albuterol sulfate (Accurate Chemical, Westbury, NY) and 0.021 g of poloxamer 188 NF grade (BASF, Mount Diive, NJ).

Solution B: A fluorocarbon-in-water emulsion stabilized by phospholipid was prepared in the following manner. The phospholipid, 1g EPC-100-3 (Lipoid KG, Ludwigshafen, Germany), was homogenized in 150g of hot deignized water (T = 50 to 60°C) using an Ultra-Turrax mixer (model T-25) at 8000 rpm for 2 to 5 minutes (T = 60-70°C). 25g of perfluorooctyl bromide (Atochem, Paris, France) was added dropwise during mixing. After the fluorocarbon was added, the emulsion was mixed for a period of not less than 4 minutes. The resulting coarse emulsion was then passed through a high pressure homogenizer (Avestin, Ottawa, Canada) at 18,000 psi for 5 passes.

Solutions A and B we're combined and fed into the spray-dryar under the conditions described above. A free flowing, white provide twas collected at the cyclone separator. The hollow procus abuteral sulfate particles had a volume-weighted mean serodynamic diameter of 1.18 ± 1.42 µm as determined by a time-offlight enelytical method (Aerosizer, Amherst Process Instruments, Amherst, MAI. Scanning electron microscopy · (SEM) enalysis showed the providers to be spherical and highly porous. The tap density of the provider was determined to be less than 0.1 glcm<sup>2</sup>.

This foregoing example serves to illustrate the inherent diversity of the present invention as a drug delivery pletform capable of effectively incorporating any one of a number of pharmaceutical agents. The principle is further illustrated in the next example.

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ΧI

#### Preparation of Hollow Perous Particles of BDP by Spray-Drying

Perforated microstructures comprising beclomethesone dipropionate (BDP) perticles were prepared by a spry-drying technique with a B-191 Mini Spray-Drier (Blüch, Flevol, Switzerland) under the following spray conditions: espiration: 100%, inlet temperature: 85°C; outlet temperature: 61°C; feed pump: 10%; N<sub>2</sub>, flow: 2,800 L/hr. The feed stock wes prepared by mixing 0.11g of lactors with a fleeroceathon-in-weter emulsion immediately prior to spray drying. The emulsion was prepared by the technique described below.

74 mg of BDP (Sigma, Chemical Co., St. Louis, MO), 0.5g of EPC-100.3 (Lipid KG, Ludwigshelan, Germany), 15mg sodium deats (Sigme), and 7mg of poloxemer 188 (BASF, Mount Olive, NJ) were dissolved in 2 ml of hot methanol. The methand was then evaporated to obtain a thin film of the phospholipidIstaroid mixture. The phospholipidIstaroid mixture was than dispersed in 64g of hot deionized water (T - 50 to 60°C) using an Ultra-Turrax mixer (model T-25) at 8000 rpm for 2 to 5 minutes (T - 60-70°C). 8 g of perfluthon (Atochem, Paris, France) was added dropvise during mixing. After the addition was complete, the emulsion was mixed for an additional period of not less than 4 minutes. The resulting coarse emulsion was than used to high pressure homogenizer (Avestin, Ottowa, Cameda) at 18,000 pil for 5 passes. This emulsion was than used to form the feed stock which was spray dried as described above. A free flowing, white powder was cellected at the cyclone separator. The hollow porcus BDP earticles had a tea density of less than 0.1 olem?

XII

#### Preparation of Hollow Porous Particles of Cromolyn Sodium by Spray-Drying

Perforated microstructures comprising cormolyn sodium were prepared by a spray-drying technique with a B-191 Mini Spray-Drier (Büchi, Flavel), Switzerland under the following spray conditions: aspiration: 100%, inlet temperature: 85°C; outlet temperature: 61°C; feed pump; 10%; N, flow: 2,800 Llhr. The feed solution was prepared by mixing two solutions A and B immediately prior to spray drying.

Solution A; 20g of water was used to dissalve 1g of cromolyn sodium (Sigma Chemical Co, St. Louis, MO) and 0.021 g of poloxemer 188 NF grade (BASF, Mount Olive, NJ).

Solution B: A fluorocarbon-in-water emulsion stabilized by phospholipid was prepared in the following manner. The phospholipid, Ig EPC-100-3 (Lipoid KG, Ludwigshafen, Germany), was homogenized in 150g of hot deenized water (T = 50 fo 60°C) using an Ultra-Turnax mixar (model T-28) at 8000 rpm for 2 to 5 minutes (T = 60-70°C). 27g of perfluorodecini (Air Products, Allentown, PA) was added dropwise during mixing. After the fluorocarbon was added, the emulsion was mixed for at least 4 minutes. The resulting coarse emulsion was then passed through a high pressure homogenizer (Avestin, Ottawa, Canada) at 18,000 psi for 5 nasses.

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Solutions A and B were combined and fed into the spray dryer under the conditions described above. A free flowing, pale yellow powder was collected at the cyclone separator. The hollow porous cromolyn sodium particles had a volume-weighted mean aerodynamic diameter of 1.23 ± 1.31 µm as determined by a time-of-flight analytical method (Aerosizer, Amharst Process Instrumenta, Amherst, MAI. As shown in Fig. 3, scanning electron microscopy (SEM) analysis showed the powders to be both hollow and porous. The tap density of the powder was determined to be less than 0.1 gicns.

#### XIII

#### Preparation of Hollow Porous Particles of DNase I by Spray-Drying

Hollow percus ONese I particles were prepared by a spray drying technique with a 8-191 Mini Spray-Drier (Büchi, Flavvil, Switzerland) under the following conditions: aspiration: 100%, inlat temperature: 80°C; outlat temperature: 61°C; feed pump; 10%; N, flow: 2,800 Libr. The feed was prepared by mixing two solutions A and B immediately prior to spray drying.

Solution A: 20g of water was used to dissolve 0.5gr of human pancreas DNase I (Calbiochem, San Diano CA1 and 0.0120 of poloxamer 188 NF orada (BASF, Mount Olive, NJI.

Solution B: A fluorocarbon-in-water emulsion stabilized by phospholipid was prepared in the following way. The phospholipid, 0.52g EPC-100 3 (Lipida KG, Ludwigshafen, Garmany), was homoganized in 87g of the dieinized water (T = 50 to 60°C) using an Ultra-Turrax mixer (model T-25) at 8000 rpm for 2 to 5 minutes (T = 60-70°C). 13g of perflubran (Atochem, Paris, France) was added dropwise during mixing. After the fluorocarbon was added, the emulsion was mixed for at least 4 minutes. The resulting course emulsion was then passed through a high pressure homoganizer (Avestin, Ottowa, Camada at 18,000 psi for 5 passes.

Solutions A and B were combined and fed into the spray dryar under the conditions described above. A free flowing, pele yellow powder was collected at the cyclone separator. The hollow porous O(8ase 1) particles had a volume-weighted mean aerodynamic diameter of  $1.29 \pm 1.40 \ \mu m$  as detarmined by a time-of-flight enalytical method (Aarosizer, Amherst Process Instruments, Amherst, MA). Scanning electron

microscopy (SEM) analysis showed the powders to be both hollow and porous. The tep density of the powder was determined to be less than 0.1 ofcm<sup>2</sup>.

The foregoing example further illustrates the extraordinary compatibility of the present invention with a variety of bioactive agents. That is, in addition to relatively small, hardy compounds such as steroids, the preparations of the present invention may be formulated to effectively incorporate larger, fragilia molecules such as contains and penaltic material.

#### XIV

#### Preparation of Perforated Ink Polymeric Particles by Spray Orying.

In the following hypothetical example, finely-divided porous spherical resin particles which may contain coloning material such as a pigment, a dye, etc. are formed using the following formulation in accordance with the teachings herein:

	Formulation:				
15	Butadiene	7.5 g		co-monor	ner
	Styrene	2.5 g		co-monor	ner
	Water	18.0 g		carrier	
	Fatty Acid Spap	0.5 g		emulsifie	r
	n-Dodecvi Mercaptan	0.050 g	modifier		
20	potassium persulfate	0.030	g	initiator	
	carbon Black	0.50 a	•		pigment

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The reaction is allowed to proceed at 50°C for 8 hours. The reaction is then terminated by spray drying the emulsion using a high pressure liquid chromatography (HPLC) pump. The emulsion is pumped through a 200 x 0.030 inch i.d. stainless steel tubing into a Nino atomizer portable spray dryer (Nino Atomize, Copenhagen, DemantAl euriped with a two fluid nozife (0.01° i.d.) employing the following settings:

Hot air flow rate:	39.5 CFM
Inlat air tamp.:	180°C
Outlet air temperature:	80°C
Atomizer nitrogen flow:	45 L/min, 1,800 psi
Liquid feed rate:	33 mL/min

It will be appreciated that unreacted monomers serve as blowing agents, creating the perforated microstructure. The described formulation and conditions yield free flowing porous polymeric particles ranging from 0.1-100, m that may be used in ind formulations. In accordance with the teachings herein the microparticles have the adventage of incorporating the pigment directly into the polymeric matrix. The process allows for the production of different particle sizes by modifying the components and the spray drying conditions with the pigment particle diameter largely dictated by the diameter of the copolymer resin particles. 5

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#### Andersen Impactor Test for Assessing MDI and DPI Performance

The MDIs and DPIs were tested using commonly accepted phermaceutical procedures. The method utilized was compliant with the United State Phermacopies (USP) procedure (Phermacopiesi Previews (1995) 22-3065-3088) incorporated herein by reference. After 5 shots to woste, 20 shots from the test MDI were made into an Andersen Impactor. The number of shots employed for assessing the DPI formulations was dictature by the drug concentration and ranged from 10 to 20 sctuations.

Extraction procedure. The extraction from all the plates, induction port, and actuator were performed in closed wide with 10 mL of a suitable solvent. The filter was installed but not assayed, because the polyecyfic binder interferred with the analysis. The mass balance and particle size distribution trends indicated that the deposition on the filter was negligibly small. Mathamol was used for extraction of becomethasone dipropianate. Delanized water was used for albutaral suffate, and cromolyn sodium. For albutaral MDIs, 0.5 ml of 1 N sodium bydroxide was added to the plate extract, which was used to convert the albutard into the phanelate form.

Quantitation procedure. All drugs were quantitated by absorption spectroscopy (Beckman DU640 spectrophotometer) relative to an external standard curve with the extraction solvent as the blank. Bedomethesone dipropionate was quantitated by measuring the absorption of the plate extracts at 23B nm Albuterol MDIs were quantified by measuring the absorption of the extracts at 243 nm, while cremelyn sodium was quantitated using the absorption peak at 326 nm.

Calculation procedure. For each MDI, the mass of the drug in the stem (component -3), actuator (2), induction port (-1) and plates (0-7) were quantified as described above. Stages -3 and -2 were not quantified for the DPI since this device was only a pirototype. The main interest was to assess the secrotypenic properties of the powder which leaves this device. The Fine Perticle Doss and Fine Particle Frestion was calculated according to the USP method referenced above. Threat deposition was defined as the mass of drug found in the induction port and on plates 0 and 1. The mean mass aerodynamic diameters (ISDI) were evaluated by fitting the experimental cumulative function with log-normal distribution by using two-parameter fitting routine. The results of these experiments are presented in subsequent examples.

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ΧVI

### Preparation of Metered Dose Inhalers Containing Hollow Porous Particles

A pre-weighed amount of the hollow porous perticles prepared in Examples I, X, XI, and XII were placed into 10 ml aluminum cans, and dried in a vacuum oven under the flow of nitrogen for 3 - 4 hours at 40°C. The amount of powder filled into the can was determined by the amount of drug required for thereneutic effect. After this, the can was crimp seeled using a DF31f50act 50 1 valve (Valois of America,

Greanwich, CT) and filled with HFA-134a (DuPont, Wilmington, DE) propallant by overpressura through the stem. The emount of the propellant in the can was determined by weighing the can before and after the fill.

XVII

Effect of Powder Porosity on MDI Performance

In order to exemine the effect powder porosity has upon the suspension stability and aerodynamic diamater, MDIs were prepared as in Exampla XIV with various preparations of perforated microstructures compaising gentamicin formulations as described in Example I. MDIs containing 0.48 wt % sprey dried powders in HFA 134a were studied. As set forth in Exemple I, the sprey dried powders exhibit varying sonsity. The formulations wern filled in clear class visits to allow for visual examination.

A strong dependence of the suspension stability and meen valume weighted serodynemic diameter was observed as a function of PFC/PC ratio and/or porosity. The valume weighted mean aerodynamic diameter (VIMAD) decreased and suspension stability increased with increasing porosity. The powders that especial solid and smooth by SEM and TEM techniques had the worst suspension stability and largest meen serodynamic diameter. MDIs which were formulated with highly porous and hollow perforated microstructures had the greatest resistance to creening and the smellest enrodynamic diameters. The measured VIMAD values for the dry opportunity of committees the properties of the dry opportunity of the properties of the properties of the dry opportunity of the properties of the dry opportunity of the properties o

Table III

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PFC/PC	Powder VMAD, $\mu$ m
0	6.1
1.1	5.9
2.2	6.4
4.8	3.9
18.8	2.6
44.7	1.8

XVIII

Comparison of Creaming Rates in Cromolyn Sodium Formulations

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A compenison of the creaming rates of the commercial Intal formulation (Rhone-Poulenc Rorer) and spray-drind hollow porous particles formulated in HFA-134a according to Example XII (i.e. see Fig. 3) is shown in Figures. 4A to 40. In each of the pictures, taken at 0 seconds, 30 seconds, 60 seconds end two hours after shaking, the commercial formulation is on the left and the perforstal microstructure dispersion formed accordance with the present invention is on the right. Whereas the commercial Intel formulation shows creaming within 30 seconds of mixing, almost no creaming is noted in the spray-dried perioles after 2 hours. Moreover, there was little creaming in perforated microstructure formulation after 4 hours (not shown). This

exempla clearly illustratas tha balanca in dansity which can be achieved when the hollow porous particles ara filled with the suspension medium (i.e. in the formation of a homodispersion).

## XIX Andersen Cascade Impactor Results for Cromolyn Sodium MDI Formulations

The results of cascede impactor tests for a commercially available product (Intal<sup>®</sup>, Rhone-Pouleac Rorer) and an analogous sprey-dried hollow porous powder in HFA-134e prepared according to Examples XII and XVI are shown below in Table IV. The tests were performed using the protocol set forth in Example XV.

Table IV

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Cromolyn Sodium MDIs					
	MMAD (GSD)	Throat Daposition, µg	Fine particle fraction, %	Fine Particle Dose, g	
Intal*,CFC (n = 4) (Rhone Poulanc) 800 µg dose	4.7 ± 0.5 (1.9 ± 0.06)	629	24.3 ± 2.1	202 ± 27	
Spray dried hollow porous powder, HFA (Alliance) (n=3)	3.4 ± 0.2 (2.0 ± 0.3)	97	67.3 ± 5.5	200 ± 11	

The MOI formulated with perforated microstructures was found to have superior serosal performence compared with Intal<sup>2</sup>. At a compareble fine particle dose, the spray dried cromolyn formulations possessed a substantially higher fine particle fraction (° 67%), and significantly decreased throat deposition (6 fold, along with a smaller MMAD value. It is important to note that the effective delivery provided for by the present invention allowed for a fine particle dose that was approximately the same as the prior art commercial formulation even though the amount of perforated microstructures edministered (300 µg) was roughly a third of the Intal<sup>2</sup> dose administered (800 µg).

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#### Companison of Andersen Cascade Impactor Results for Albuteral Sulfate Microspheres Delivered From DPIs and MDIs

The in vitro aerodynamic properties of hollow porous albuterol sulfate microspheres as prepared in Exemple X was characterized using an Andersen Mark II Cascade Impactor (Andersen Sampler, Atlanta, GA) and an Amharst Aerosizer (Amherst Instruments, Amherst, MA).

<u>OP1 testing.</u> Approximately, 300mcg of spray-dried microspheres was loaded into a proprietary inhelation device. Activation and subsequent plane generation of the dry powder was achieved by the actuation of 50 µl of pressurized HFA 134s through a long induction tube. The pressurized HFA 134s forced gir through the induction tube toward the sample chamber, and subsequently sercedized a plume of dry

provider into the air. The dry powder plume was then taken in the cascade impactor by means of the air flow through drawn through the testing device. A single actuation was discharged into the aerosizer sample chamber for particle size analysis. Ten actuations were discharged from the device into the impactor. A 30 second interval was used between each actuation. The results were quantitated as described in Example XV.

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MDI testing. A MDI preparation of albuteral sulfate microspheres was prepared as in Exemple XVI.

A single actuation was discharged into the aerosizer sample chamber for particle size analysis. Twenty actuations were discharged from the device into the impactor. A 30 second interval was used between each actuation. Again, the results were quantitated as described in Exemple XVI.

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The results companing the particle size enalysis of the nest albuterol sulfate powder and the albuterol sulfate powder discharged from either a DPI or MDI are shown in Table V balow. The albuterol sulfate powder delivered from the OPI was indistinguishable from the nest powder which indicates that little or no aggregation had occurred during exclusion. On the other hand, same aggregation was obsarved using an MDI as evidenced by the larger errodynamic disnetter of particles delivered from the device.

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V eldaT

Sample	Mean Size (µm)	% under 5.4 $\mu$ m	95% under (µm)
Neat powder	1.2	100	2.0
MOI	2.4	96.0	5.1
DPI	1.1	100	1.8

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Similar results were observed when compering the two dosage forms using an Andersen Cascade impactor (Figure 5). The spra-ydried abutard suffets powder delivered from the DPI had enhanced deep lung deposition and minimized throat deposition when compared with the MDI. The MOI formulation had a fine particle fraction (FP) of 79% and a fine particle dose (FPD) of 77 µg/actuation, while the OPI had a FPF of 87% and a FPD of 100µg/ actuation.

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Figure 5 and the Example above exemplifies the excellent flow and ecrodynamic properties of the herein described spray-fixed providers delivered from a OPI. Indeed, one of the primary advantages of the present invention is the ability to produce small aerodynamically light particles which aerosolize with ease and which have excellent inhalation properties. These powders have the unique properties which enable them to be effectively and efficiently delivered from either a MDI or DPI. This principle is further illustrated in the next Example.

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Comparison of Andersen Cascade Impactor Results for Beclomethasone Dipropionata Microspheres Delivered From QPIs and MOIs

The in vitro sernolynamic properties of hollow perous becomethasons dipropinate (80P) microspheres as prepared in Example XI was characterized using an Andersen Mark II Cascade Impactor (Andersen Samuler, Atlanta, GA) and an Amberst Aerosizer (Amberst Instruments, Amberst, MA).

DPI testing. Approximately, 300µg of sprey-dried microspheres was loaded into a proprietary inhalation device. Activation and subsequent plume generation of the dry powder was achieved by the actuation of 50 µl of pressurized HFA 134a through a long induction totle. The pressurized HFA 134a forced air through the induction tube toward the sample chamber, and subsequently aerospized a plume of dry powder into the air. The dry powder plume was then taken in the cascade impactor by means of the air flow through drawn through the testing device. A single actuation was discharged into the serosizer sample chamber for particle size analysis. Twenty actuations were discharged from the device into the impactor. A 30 second interval was used between each actuation.

MDI testing. A MOI preparation of becomethesone dipropionate (8DP) microspheres was prepared as in Example XVI. A single actuation was discharged into the serosizer sample chamber for particle size analysis. Twenty actuations were discharged from the device into the impactor. A 30 second interval was used between each actuation.

The results comparing the particle size analysis of the neat BDP powder and the BDP powder discharged from either a DPI or MDI are shown in Table VI immediately below.

Table VI

Sample	Mean Size (¿/m)	% under 5.4 µm	95% under (µm)
Neat powdar	1.3	100	2.1
MDI	2.2	98.1	4.6
DPI	1.2	99.8	2.2

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As with Example XX, the BDP powder delivered from the DPI was indistinguishable from the neat powder which indicates that little or no aggregation had occurred during actuation. On the other hand, some aggregation was observed using an MDI as evidenced by the larger aerodynamic diameter of particles delivered from the davice.

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The sprsy-died BDP powder delivered from the DPI had enhanced deep lung deposition and minimized throat deposition when compared with the MDI. The MDI formulation had a fire particle fraction (FPF) of 79% and a fine particle dose (FPD) of 77.  $\mu$ pjactustion, while the DPI had a FPF of 87% and a FPD of 100 $\mu$ col actuation.

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This foregoing example serves to illustrate the inherent diversity of the present invention as a drug delivery platform capable of effectively incorporating any one of a number of pharmacounical agents and effectively delivered, from various types of delivery devices (here MDI and DPI) currently used in the

pharmaceutical arana. The axcellent flow and serodynamic properties of the dry powders shown in the proceeding exemples is further exemplified in the next example.

# XXII <u>Comparison of Andersen Cascade Impactor Results for</u> Albuterol Sulfata Microspheras and Ventolin Rotacaps from a Rotahaler Device

The following procedure was followed to compare the inhalation properties of Ventolin Rotocaps' (a commercially available formulation) vs. albutarel sulfate hollow percus microsphares formed in accordance with the present invention. Both prepartiens were discharged from a Rotohaler' device into an 8 stage Andersen Mark II cascade impactor operated at a flow of 60Llmin. Preparation of the albutarol sulfate microspheres is described in Exemple X with albutarol sulfate deposition in the cascade impactor analyzed as described in Exemple XV. Approximately 300 µg of albutarol sulfate microsphares were manually loaded into empty Ventdin Rotocap' gelatin capsules. The procedure described in the package insert for loading and actuating drug capsulas with a Rotohaler' device was followed. Ten actuations were discharged from the device into the impactor. A 30 second interval was used between each actuations.

The results comparing the cascade impactor analysis of Ventolin Rotocaps' and hollow porous albuterol sulfate microspheres discharged from a Rotohaler' device are shown in Table VI immediately below.

Teble VII

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Sample	MMAD	Fine Particle Fraction	Fine Particle Dose
	(GSD)	%	(mcg/dose)
Ventolin Roteceps (n = 2)	7.869	20	15
	(1.6064)	1	
Albuterol Sulfate	4.822	63	60
Microsobares (p = 3)	(1.9082)	1	

The hollow porous albuterol sulfate powder delivered from the Rotohaler device had a significantly higher fine perticle fraction CF-India and a smaller MMAD value as compared with Ventolin Rotocapt. In this regard, the commercially available Ventolin Rotocap\* formulation had a fine particle fraction (FPF) of 20% and a fine particle dose (FPD) of 15 pylactuation, whereas the hollow porous albuterol sulfate microspheres had a FPF or 63% and a FPD of 800zal actuation.

The example above exemplifies the excellent flow and serodynamic properties of the spray-dried powders delivered from a Rotahaler device. Moreover, this example demonstrates that fine powders can be effectively delivered without carrier particles.

XXIII

<u>Nebulization of Porous Perticulate Structures Comprising</u>

Phospholipids and Cromolyn sodium in Perfluorooctylethana

#### using a MicroMist Nebulizer

Forty milligrams of the lipid based micrespheres containing 50% cromolyn accidium by weight (as time Example XII) were dispersed in 10 rdl perfluoroactylethane (PFOEI) by shaking, forming a suspension. The suspension was nebulized until the fluorocarhon liquid was delinered or had evaporated using a MicroMist (DeVillissa) dispeasable impactor was used to measure the resulting particle size distribution. More specifically, cromolyn acidium content was measured by UV adsorption at 326mm. The fine particle fraction is the ratio of particles deposited in stages 2 through 7 to those deposited in all stages of the impactor. The fine particle mass is the weight of material deposited in stages 2 through 7. The deep lung fraction is the retio of particles deposited in stages 5 through 7 the impactor (which correlate to that sivedii) to those deposited in all stages. The deep lung mass is the weight of material denosited in states 5 through 7. The day VIII immediately below oranges assummany of the results.

Tehlo VIII

Fina particla fraction	fine particle mass	deep lung fraction	deap lung mass
90%	6 mg	75%	5 mg

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#### XXIV

# Nebulization of Porous Particulate Structures Comprising Phospholicids and Cromolyn Sodium in Perfluorocctylethene

#### using a Raindrop® Nebulizar

A quantity of lipid based microspheres containing 50% cromolyn sodium, as from Exemple XII, weighing 40 mg was dispersed in 10 ml perfluoreoctylethane (PFOE) by shaking, thereby forming a suspension. The suspension was nebitined until the fluoreocathon liquid was oldered or had evapersed using a Raindrop disposable nebuliter (Nidloor Putten Bennett connected to a PulmoAide" air compressor (DeVilbias). An Andersen Cascade Impuctor was used to measure the residing particle size distribution in the manner described in Examples XV and XXIII. Teble IX immediately below provides a summer of the needs.

Table IX

Fine particle fraction	fine particla mass	Deep lung fraction	deep lung mass
90%	4 mg	80%	3 mg

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The contents of plastic vial conteiting a unit dose inhalation adultion of 20 mg of cromolyn sodium in 2 ml purified water (Dey Laboratories! was rehulized using a MicroNict disposable nebulizer (De/Nicks using a Putmahide<sup>2</sup> air compressor (De/Nicks). The cromelyn sodium sodium nebulized for 30 minutes. An Andersen Cescade Impactor was used to measure the resulting size distribution of the nebulized particles, by the method described shown in Exemple XV. Table X immediately below provides a summer of the results.

Table X

fine particle fraction	fina particla mass	Deap lung fraction	Deep lung mass
90%	7 mg	60%	5 mg

With regard to the instant results, it will be appreciated that, the formulations nebulized from fluorecarbon suspension mediums in Exemples XXIII and XXIV provided a greater percentage of deep lung deposition than the suspenses solution. Such high deposition rates deep in the lung is particularly desirable when delivering agents to the systemic circulation of a sedient.

Those skilled in the art will further appreciate that the present invention may be emboded in other specific forms without departing from the spirit or central attributes thereof. In that the foregoing description of the present immention discloses only exemplary embodiments thereof, it is to be understood that to ther variations are contemplated as being within the scope of the present invention. Accordingly, the present invention is not limited to the particular embodiments which have been described in detail herein. Rather, reference should be made to the appreciacion as sindective of the scope and content of the invention.

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#### WHAT IS CLAIMED:

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Use of a bioscrive agent in the manufacture of a medicament for pulmonary delivery whereby
the medicament comprises a plurality of perforated microstructures which are sensolized using an inhalation device
to provide sensoslized medicament comprising said bioactive agent wherein said sensolized medicament is in a form
for administration to at least a portion of the nasal or pulmonary air passages of a patient in need thereof.

- The use of claim 1 wherein said inhalation device comprises a metered dose inhalar, a dry powder inhalar or a nebutizer.
  - The use of claim 1 wherein said parforated microstructures are in the form of a dry powder.
- The use of claim 1 wherein said perforated microstructures are dispersed in a nonaqueous suspension medium.
  - The use of any of claims 1 to 4 wherein said perforated microstructures comprise a surfactant.
- The use of claim 5 wherein said surfactant is selected from the group consisting of phospholipids, nonionic detergents, nonionic block copolymers, ionic surfactants, biocompetible fluorinated surfactants and combinations thereof.
  - 7. The use of claims 5 or 6 wherein said surfactant is a phospholipid.
- B. The use of claim 7 wherein said phospholipid is selected from the group consisting of disurcylphosphatidylcholine, dispelmitolyphosphatidylcholine, disteroylphosphatidylcholine, disteroylphosphatidylcholine, disteroylphosphatidylcholine, disteroylphosphatidylcholine and combinations thereof.
- 9. The use of any of claims 1 to 8 wharein the mean aerodynamic diameter of the perforated microstructures is between 0.5 and 5  $\mu$ m.
- The use of any of claims 1 to 9 wherein said perforated microstructures have a bulk density of less than about 0.5 g/cm².
- The use of any of claims 1 to 10 wherein said perforated microstructures have a mean geometric diameter of less than about 6 µm.
  - 12. The use of any of claims 1 to 11 wherein said bioactive agent is selected from the group consisting of entail enrice, bronchodilators, pulmonery lung surfactants, analysicis, ambibiotics, leukstriene inhibitors or entagonists, andisistemines, entitinflamentories, entimeoplastics, enticholinenzics, entertuiscs, anti-tuberculars, imaging agents, cardiovascular agents, enzymes, storeids, generic meterial, viral vectors, entiennes agents, proteins, conditions and productive and combinations thereof.
    - A method for forming a perforated microstructure comprising the steps of: providing a liquid feed stock comprising on active agent; atomizing said flouid feed stock to produce dispessed liquid droplets;

drying said liquid droplets under predetermined conditions to form perforeted microstructures comprising said active exent and

collecting said perforated microstructures.

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- The method of claim 13 wherein said feed stock comprises a blowing agent.
- 15. The method of claim 14 wherein said blowing agent comprises a nonfluorinated oil.
  - 16. The method of claim 14 wherein said blowing agent comprises a fluorinated compound.
- The method of claim 16 wherein said fluorinated blowing agent has a builing point greater than about 60°C.
- 18. The method of any of claims 13 to 17 wherein said feed stock comprises a colloidal system.
- 19. The method of any of claims 13 to 18 wherein said feed stock comprises a surfactant.
- 20. The method of claim 19 wherein said surfactant is selected from the group consisting of phospholipids, nonionic datargents, nonionic block copplymers, ionic surfactants, biocompatible fluorineted surfactants and combinations thereof.
  - 21. The method of claim 19 or 20 wherein said surfactant is a phospholipid.
- 2. The method of claim 21 wherein said phospholipid is selected from the group consisting of dilaury (phosphetidylcholine, diohyphosphetidylcholine, dipalmit vylphosphetidylcholine, distartolylchosphetidylcholine, distartolylcholine, distartolylchosphetidylcholine, distartolylcholine, distartolylchosphetidylcholine, distartolylcholine, distartolylc
- The method of any of claims 13 to 22 wherein said collected perforated microstructures comprise hollow perous microsphares.
- 25. The method of any of claims 13 to 24 wherein said perforated microstructuras have a mean geometric diameter of less than about 5 µm.
- The method of any of claims 13 to 25 wherein said active agent comprises a bioactive agent.
  - 27. The method claim 26 wherein said bisective egent is selected from the group consisting of antiallates, bronchodilates, pulmonery lung surfactants, analgasics, antibibites, leukoninee inhibiters or antegonists, antibistanzines, entitindiammatories, entineoplastics, anticholinergics, anesthebics, enti-tuberciders, imaging agents, cardiovascular agents, enzymes, steroids, genetic meterial, viral vectors, entisense agents, proteins, esandées and combinations themed.
  - 28. The method of any of claims 13 to 27 wherein said atomization step is accomplished using a soray dryer.
    - 29. A perforated microstructure formad according to any one of claims 13 to 28.
- 35 30. A method for increasing the dispersibility of a powder comprising the steps of:

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providing a liquid feed stock comprising an active agent; and

sprey dying said liquid feed stock to produce a perforated microstructure powder having a bulk density of less than about 0.5 g/cm<sup>2</sup> wherein said powder exhibits reduced van der Wasls attractive forces when commend to a relatively non-corous powder of the same composition.

- 31. The method of claim 30 wherein said liquid feed stock comprises a blowing agent
  - 32. The mathod of claim 31 wherein said blowing agent comprises a nonfluorinated oil.
  - 33. The method of claim 31 wherein said blowing agent comprises a fluorinated compound.
  - 34. The method of claim 33 wherein said fluorinated compound has a boiling point of greater than about 60°C.
    - 35. The method of any of claims 30 to 34 wherein said feed stock comprises a surfactant.
  - 38. The method of claim 35 wherein said surfactant is selected from the group consisting of phospholipids, nonionic detargents, nonionic block copolymers, ionic surfactants, biocompatible fluorinated surfactants and combinations thereof.
    - 37. The method of claim 35 or 36 wherein said surfectant is a phospholipid.
  - 38. The method of claim 37 wherein said phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of dilaur of phospholipid is selected from the group consisting of the group consis
  - 39. The method of any of claims 30 to 38 wherein said perforated microstructures comprise hollow porous microspheres.
    - 40. The method of any of claims 30 to 39 wherein said active agent comprises a bioactive agent.
  - 41. The method daim 40 wherein said bisective agent is selected from the group consisting of entiellaritys, brancho diletors, pulmonary lung surfactants, endejestes, antibiotics, leukutriene inhibitors or antigonists, entitistamines, entiinflammentaries, antienglastics, entitholinergics, anasthetics, anti-tuberculars, imaging agents, cardivactular agents, enzymes, steroids, genetic material, viral vectors, antisense agents, propriess, peptides and combinations thereof.
    - 42. A perforated microstructure powder formed according to any one of claims 30 to 41.
  - 43. A powder having increased dispersibility comprising a plurality of perforated microstructures having a bulk density of less than about 0.5 glcm² wherein said perforated microstructure powder comprises an active seent.
    - 44. The powder of claim 43 wherein said powder comprises hollow porous microspheres.
  - 45. The powder of claims 43 or 44 wherein the mean aerodynamic diameter of said perforated microstructures is between 0.5 and 5 µm.
- 46. The powder of any of claims 43 to 45 wherein said perforated microstructures have a 35 mean geometric diemeter of less than about 5 µm.

 The powder of any of claims 43 to 46 wherein said perforated microstructures comprise a surfactant.

- 48. The provder of claim 47 wherein said surfactant is selected from the group consisting of phospholipids, nonionic detergents, nonionic block copolymers, ionic surfactants, biocompatible fluorinated surfactants and combinations thereof.
  - 49 The nowder of claim 47 or 48 wherein said surfactant is a phospholipid.

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- 50. The powder of claim 49 wherein said phospholipid is selected from the group consisting of disuncythosphatidylcholine, diologiphosphatidylcholina, dipalmitorychosphatidylcholine, distarroythosphatidylcholine dibahenoyghosphatidylcholine, distarchidovychosphatidylcholine and combinations thereof.
  - 51. The powder of claim any of claims 43 to 50 wherein said active agent is a bioactive agent.
- 52. The powder of claim 51 wherein said bioscive agent is selected from the group consisting of antiallargics, bronchodilators, pulmonary lung surfactants, analyssics, antibiotics, laukotriens inhibitors or antagonists, antibistamines, antiinflammatories, antineoplastics, antibiotics, laukotriens inhibitors or antagonists, antibistamines, antiinflammatories, antineoplastics, antibiotics, laukotriens, antistamines, antiinflammatories, antipides, anticologistics, anticologisti
- An inhalation system for the pulmonary administration of a bioactive agent to a patient comprising:

an inhalation device comprising a reservoir; and

- a powder in said reservoir wherein said powder comprises a plurality of parforated microstructures having a bulk density of less than about 0.5 glcm<sup>3</sup> wherein said perforated microstructure powder comprises a bloactive agent whereby said inhalation device provides for the aerosolized administration of said powder to at least a portion of the nasal or pulmonary air passages of a patient in need thereof.
- 54. The system of claim 53 wherein said inhalation device comprises a dry provider inhaler, a metered dose inhaler or a nebulizer.
- 55. The system of claim 53 wherein said perforated microstructures are dispersed in a nonaqueous suspension medium.
- The system of claim 55 wherein said nonequeous suspension medium comprises a fluorinated compound.
  - 57. The system of any of claims 54 to 56 wherein said perforated microstructures comprise a surfactant.
  - 58. The system of claim 57 Wherein said surfactant is selected from the group consisting of phospholipids, nonionic detergents, nonionic block copolymers, ionic surfactants, biocompetible fluorinated surfactants and combinations thereof.

59. The system of claims 57 or 58 wherein said surfactant is a phospholipid.

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- The system of any of claims 54 to 59 wherein said bisective agent is selected from the group consisting of antiallergics, transcholistors, pulmosny lung surfactants, an anlegacis, antibiotics, laukatione inhibitors or entegorists, antibisterarizes, antiintermetories, antimeoplastics, anticholinergics, anesthetics, enti-tuberculars, imaging agents, cardiovascular agents, enzymes, stereids, genetic metorial, viral vectors, antisense agents, proteins, anotides and comminations theme?
- 61. A method for the pulmonary delivery of one or more biosocitive agents comprising the steps of: providing a powder comprising a plurality of perforated microstructures having a bulk density of less than about 0.5 a/cm² wherein said perforated microstructure powder comprises a bioactive agent;

ecrosolizing said perforated microstructure powder to provide an aerosolized medicament; and administering a thempeutically affective encount of said serosolized medicament to at least a portion of the nassal or pulmonary air passages of a patient in need thereof.

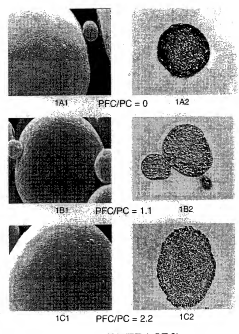


FIG. 1 (SHEET 1 OF 2)

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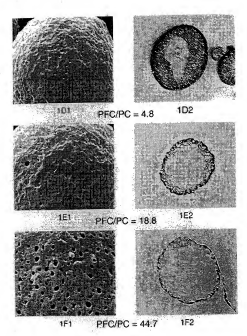
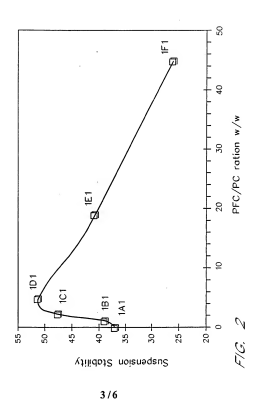


FIG. 1 (SHEET 2 OF 2)

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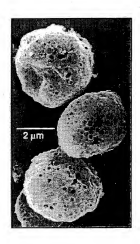
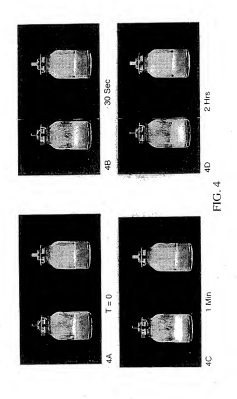
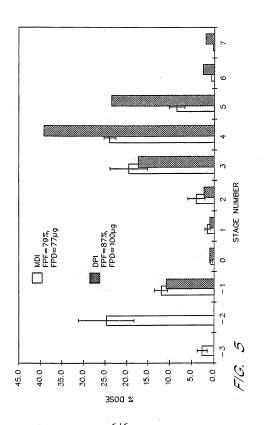


FIG.3

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b .ational Application N

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A. CLASSII IPC 6	FICATION OF SUBJECT MATTER A61K9/00 A61K9/51								
According to	International Petent Classification (IPC) or to both national classifica	tion and IPC							
B. FIELDS SEARCHED									
Minimum documentetion searched (classification system tollowed by classification symbols)									
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*** document which may throw doubts on priority claimly) or which is clied to establish the publication date of another which is clied to establish the publication date of another checklon or other special reason (as specified).  **Comment where he not not decisionar, use, a withfullion or document to consider to browthe an inventible step when the consideration and or other dates to decisionary.									
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Name and r	meling eddress of the ISA European Patient Office, P.B. 5818 Patentiaan 2 N 2280 HV Rijswijk Tel. (431-70) 340-2040, Tr. 31 651 epo nl. Farr (431-70) 340-3010	Authorized officer Fischer	, w						

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